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(54) Title: RECOMBINANT HERPESVIRUS OF TURKEYS AND USES THEREOF

#### (57) Abstract

This invention provides a recombinant herpesvirus of turkeys comprising a foreign DNA sequence encoding a cytokine inserted into an insertion region which comprises a XhoI site within an EcoR1 #9 fragment of a herpesvirus of turkeys viral genome, and the foreign DNA sequence encoding a cytokine which is capable of being expressed in a host cell infected with the herpesvirus of turkeys. This invention provides a recombinant herpesvirus of turkeys-Marek's disease virus chimera comprising a herpesvirus of turkeys unique long viral genome region and a Marek's disease virus unique short region. Lastly, this invention provides homology vectors for producing a recombinant herpesvirus of turkeys, host cells, and vaccines and methods for immunization.

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### RECOMBINANT HERPESVIRUS OF TURKEYS AND USES THEREOF

This application is a continuation of U.S. Serial No. 08/362,240, filed December 22, 1994, which is a continuation-in-part of 08/288,065, filed August 9, 1994, the contents of which are hereby incorporated by reference into.

Throughout this application various publications are referenced by Arabic numerals in parenthesis. Full citations for these publications may be found at the end of the specification immediately preceding the claims. The disclosures of these publications are in their entirety hereby incorporated by reference into this application to more fully describe the state of the art to which this invention pertains.

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#### BACKGROUND OF THE INVENTION

The ability to isolate DNA and clone such isolated DNA into bacterial plasmids has greatly expanded the approaches available to make viral vaccines. The methods used to make the present invention involve modifying cloned DNA sequences from various viral pathogens of animals, by insertions, deletions, single or multiple base changes, and subsequent insertions of these modified sequences into the genome of the virus. One utility of the addition of a foreign sequence is achieved when the foreign sequence encodes a foreign protein that is expressed during viral infection of the animal. The resulting live virus may then be used in a vaccine to elicit an immune response in a host animal

and provide protection to the animal against disease. A virus with these characteristics is referred to as a viral vector, because it becomes a living vector that will carry and express the foreign protein in the host animal. In effect it becomes an elaborate delivery system for the foreign protein(s).

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More specifically, the present invention relates to the use of herpesvirus of turkeys (HVT) as a viral vector for vaccination of birds against disease. The group of herpesviruses comprise various pathogenic agents that infect and cause disease in a number of target species:

swine, cattle, chickens, horses, dogs, cats, etc. Each herpesvirus is specific for its host species, but they are all related in the structure of their genomes, their mode of replication, and to some extent in the pathology they cause in the host animal and in the mechanism of the host immune response to the virus infection.

The application of recombinant DNA techniques to animal 25 viruses has a relatively recent history. The first viruses to be engineered have been those with the smallest genomes. In the case of the papovaviruses, because these viruses are so small and cannot accommodate much extra DNA, their use in genetic 30 engineering has been as defective replicons. gene expression from these viruses requires a wild-type helper virus and is limited to cell culture systems. For adenoviruses, there is а small amount of nonessential DNA that can be replaced by foreign 35 sequences. The only foreign DNA that seems to have

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been expressed in adenoviruses are the T-antigen genes from papovaviruses (Mansour, et al., Proc. Natl. Acad. Sci. US, 1985; Thummel, et al., Cell, 1983; Scolnick, et al., Cell, 1981; Thummel, et al., Cell, 1981), and the herpes simplex virus (HSV) thymidine kinase gene (Haj-Ahmed and Graham, J. of Virology, 1986). These publications do not identify the nonessential regions in HVT wherein foreign DNA may be inserted, nor do they teach how to achieve the expression of foreign genes in HVT, e.g., which promoter sequence and termination sequence to use.

15 Another group of viruses that have been engineered are the poxviruses. One member of this group, vaccinia, has been the subject of much research on foreign gene Poxviruses are large DNA-containing expression. viruses that replicate in the cytoplasm of the infected 20 cell. They have a structure that is unique in that they do not contain any capsid that is based upon symmetry or helical icosahedral symmetry. poxviruses are most likely to have evolved from bacterial-like microorganisms through the loss of In part due to this 25 function and degeneration. advances made in the genetic uniqueness, the engineering of poxviruses cannot be directly extrapolated to other viral systems, including herpesviruses and HVT. Vaccinia recombinant virus 30 constructs have been made in a number of laboratories that express the following inserted foreign genes: HSV thymidine kinase gene (Mackett, et al., Proc. Natl. Acad. Sci. USA, 1982; Panicali and Paoletti, Proc. Natl. Acad. Sci. USA, 1982, hepatitis B surface antigen (Paoletti, et al., Proc. Natl. Acad. Sci. USA, 1984; 35

Smith et al., Nature, 1983), HSV glycoprotein D gene, influenzae hemagglutinin gene (Panicali, et al., Proc. Natl. Acad. Sci. USA, 1983; Smith, et al., Proc. Natl. Acad. Sci. USA, 1983), malaria antigen gene (Smith, et Science. 1984, and vesicular stomatitis glycoprotein G gent (Mackett, et al., Science, 1986). The general overall features of vaccinia recombinant DNA work are similar to the techniques used for all the viruses, especially as they relate to the techniques in reference (Maniatis, et al., Molecular Cloning, 1982). However in detail, the vaccinia techniques are not applicable to herpesviruses and HVT. The utility of vaccinia as a vaccine vector is in question because of its close relationship to human smallpox and its known pathogenicity to humans. Thus, the use of the hostspecific herpesvirus HVT is a better solution to vaccination of poultry.

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Among the primate herpesviruses, only HSV of humans and, to a limited extent, herpes saimiri of monkeys have been engineered to contain foreign DNA sequences. The first use of recombinant DNA to manipulate HSV involved cloning a piece of DNA from the L-S junction region into the unique long region of HSV DNA, specifically into the thymidine kinase gene (Moccarski, et al., Cell, 1980). This insert was not a foreign piece of DNA, rather it was a naturally occurring piece of herpesvirus DNA that was duplicated at another place in the genome. This piece of DNA was not engineered to specifically express a protein, and thus this work does not involve expression of protein in herpesviruses. The next manipulation of HSV involved the creation of deletions in the virus genome by a combination of

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recombinant DNA techniques and thymidine kinase selection. Using this approach, the HSV alpha-22 gene has been deleted (Post, et al., Cell, 1981), and a 15,000 basepair sequence of DNA has been deleted from the internal repeat of HSV (Poffenberger, et al., Proc. Natl. Acad. Sci. USA, 1981).

The following cases involve insertion of genes that 10 encode protein into herpesviruses: the insertion of HSV glycoprotein C into a naturally occurring deletion mutant of this gene in HSV (Gibson and Spear, J. of Virology, 1983); the insertion of glycoprotein D of HSV type 2 into HSV type 1 (Lee, et al., Proc. Natl. Acad. 15 Sci. USA, 1982), with no manipulation of promoter since the gene is not 'foreign'; the sequences insertion of hepatitis B surface antigen into HSV under the control of the HSV ICP4 promoter (Shih, et al., Proc. Natl. Acad. Sci. USA, 1984); and the insertion of 20 bovine growth hormone into herpes saimiri virus with an SV40 promoter (the promoter did not work in this system and an endogenous upstream promoter served to transcribe the gene) (Desrosiers, et al., 1984). 25 additional foreign genes (chicken ovalbumin gene and Epstein-Barr virus nuclear antigen) have been inserted into HSV (Arsenakis and Roizman. 1984). and glycoprotein X of pseudorabies virus has been inserted into HSV (Post, et al., 1985).

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These cases of deletion or insertion of genes into herpesviruses demonstrate that it is possible to genetically engineer herpesvirus genomes by recombinant DNA techniques. The methods that have been used to

insert genes involve homologous recombination between the viral DNA cloned in plasmids and purified viral DNA transfected into the same animal cell. However, the extent to which one can generalize the location of the deletion and the sites for insertion of foreign genes is not known from these previous studies.

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10 One object of the present invention is a vaccine for Marek's disease. Marek's disease virus (MDV) is the causative agent of Marek's disease which encompasses fowl paralysis, a common lymphoproliferative disease of The disease occurs most commonly in young chickens between 2 and 5 months of age. 15 The prominent clinical signs are progressive paralysis of one or more of the extremities, incoordination due to paralysis of legs, drooping of the limb due to wing involvement, and a lowered head position due to involvement of the neck 20 muscles. In acute cases, severe depression may result. In the case of highly oncogenic strains, there is characteristic bursal and thymic atrophy. In addition, there are lymphoid tumors affecting the gonads, lungs, liver, spleen, kidney and thymus (Mohanty and Dutta, 25 1981).

Most chickens are vaccinated against MDV at one day of age to protect the bird against MDV for life. Prior to the present invention, the principal vaccination method for MDV involved using naturally occurring strains of turkey herpesvirus (HVT). It would be advantageous to incorporate other antigens into this vaccination at one day of age, but efforts to combine conventional vaccines have not proven satisfactory to date due to

competition and immunosuppression between pathogens. The multivalent HVT-based vaccines engineered in this invention represent a novel way to simultaneously vaccinate against a number of different pathogens. For the first time, a recombinant HVT with a foreign gene inserted into a non-essential region of the HVT genome is disclosed.

The types of genetic engineering that have been performed on these herpesviruses consist of cloning parts of the virus DNA into plasmids in bacteria, reconstructuring the virus DNA while in the cloned state so that the DNA contains deletions of certain sequences, and furthermore adding foreign DNA sequences either in place of the deletions or at sites removed from the deletions.

A foreign gene of interest targeted for insertion into the genome of HVT may be obtained from any pathogenic organism of interest. Typically, the gene of interest will be derived from pathogens that in poultry cause diseases that have an economic impact on the poultry industry. The genes may be derived from organisms for which there are existing vaccines, and because of the novel advantages of the vectoring technology the HVT derived vaccines will be superior. Also, the gene of interest may be derived from pathogens for which there is currently no vaccine but where there is a requirement for control of the disease. Typically, the gene of interest encodes immunogenic polypeptides of the pathogen, and may represent surface proteins, secreted proteins and structural proteins.

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A relevant avian pathogen that is a target for HVT vectoring is Infectious Laryngotracheitis virus (ILTV). ILTV is a member of the herpesviridiae family, and this pathogen causes an acute disease of chickens which is characterized by respiratory depression, gasping and expectoration of bloody exudate. Viral replication is limited to cells of the respiratory tract, where in the trachea the infection gives rise to tissue erosion and hemorrhage. In chickens, no drug has been effective in reducing the degree of lesion formation or decreasing clinical signs. Vaccination of birds with various modified forms of the ILT virus derived by cell passage and/or tedious regimes of administration have conferred acceptable protection in susceptible chickens. Because of the degree of attenuation of current ILT vaccines care must be taken to assure that the correct level of virus is maintained; enough to provide protection, but not enough to cause disease in the flock.

An additional target for the HVT vectoring approach is Newcastle disease, an infectious, highly contagious and debilitating disease that is caused by the Newcastle disease virus (NDV). NDV is a single-stranded RNA virus of the paramyxovirus family. The various pathotypes of NDV (velogenic, mesogenic, lentogenic) differ with regard to the severity of the disease, the specificity and symptoms, but most types seem to infect the respiratory system and the nervous system. NDV primarily infects chickens, turkeys and other avian species. Historically vaccination has been used to prevent disease, but because of maternal antibody interferences, life-span of the bird and route of

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administration, the producer needs to adapt immunization protocols to fit specific needs.

The therapeutic agent that is delivered by a viral vector of the present invention must be a biological molecule that is a by-product of swinepox virus replication. This limits the therapeutic agent in the first analysis to either DNA, RNA, or protein. are examples of therapeutic agents from each of these classes of compounds in the form of anti-sense DNA, anti-sense RNA (S. Joshi, et al., J. of Virology, 1991), ribozymes (M. Wachsman, et al., J. of General Virology, 1989), suppressor tRNAs (R.A. Bhat, et al., Nucleic Acids Research, 1989), interferon-inducing double stranded RNA and numerous examples of protein therapeutics, from hormones, e.q., insulin, lymphokines, e.g., interferons and interleukins, to naturals opiates. The discovery of these therapeutic agents and the elucidation of their structure and function does not make obvious the ability to use them in a viral vector delivery system.

## SUMMARY OF THE INVENTION

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This invention provides a recombinant herpesvirus of turkeys comprising a foreign DNA sequence encoding a cytokine inserted into an insertion region which comprises a XhoI site within a EcoR1 #9 fragment of a herpesvirus of turkeys viral genome, and the foreign DNA sequence encoding a cytokine which is capable of being expressed in a host cell infected with the herpesvirus of turkeys.

Lastly, this invention provides homology vectors for producing a recombinant herpesvirus of turkeys, host cells, and vaccines and methods for immunization.

## BRIEF DESCRIPTION OF THE FIGURES

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# Figures 1A-1C: Details of HVT Construction and Map

Figure 1A shows BamHI restriction fragment map of the HVT genome. Fragments are numbered in order of decreasing size; letters refer to small fragments whose comparative size has not been determined.

Figure 1B shows BamHI #16 fragment of the HVT genome showing location of  $\beta$ -galactosidase gene insertion in S-HVT-001.

Figure 1C shows BamHI #19 fragment of the HVT genome showing location of  $\beta$ -galactosidase gene insertion.

Legend: B = BamHI; X = XhoI; H = HindIII; P = PstI; S = SalI; N = NdeI; R = EcoRI.

## Figures 2A-2D: Insertion in Plasmid 191-47.

Figure 2A contains a diagram showing the orientation of DNA fragments assembled in plasmid 191-47. Figures 2A to 2D show the sequences located at each of the junctions between the DNA fragments in plasmid 191-47. (SEQ ID NOs: 20, 21, 22, 23, 24, 25, 26, and 27).

## Figures 3A-3B: Details of S-HVT-003 Construction.

Figure 3A shows restriction map of HVT DNA in the region of the BamHI #16 fragment. This fragment is contained within large HindIII fragment. Figure

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3A also shows the XhoI site which was first changed to an *EcoRI* (R) site by use of a "linker" and standard cloning procedures. Figure 3A also shows details of the construction of the beta-gal gene and IBVD gene inserted into the *BamHI* #16 fragment for use in homologous recombination. Both genes were under the control of the PRV gX gene promoter (qX).

Figure 3B show the S-HVT-003 genome, including the location of the two inserted foreign genes,  $\beta$ -gal and IBDV.

In Figure 3: H = HindIII; B = BamHI; X = XhoI;

R = EcoRI; Xb = XbaI; Hp = HpaI; S = SmaI; UL = unique long region; US = unique short region; IR = internal repeat region; TR = terminal repeat region.

#### 20 Figure 4:

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Western blot indicating the differential expression of the IBDV 32kD antigen in cellular lysates of S-HVT-003 infected cells (32kD present) and S-HVT-001 infected cells (32kD negative). IBDV specific polypeptides were identified by probing the blot with hyper-immune rat antiserum directed against denatured IBDV virions. This serum reacts primarily with the immunodominant 32kD antigen (IBDV VP3). The lanes on the blot contain: 1) protein molecular weight standards, 2) uninfected CEF cells, 3) S-HVT-001 infected CEF's, 4) 5) & 6) S-HVT-003 and 7) IBDV virion polypeptides.

#### 35 Figure 5:

Western blot indicating the differential expression of the 42kD (VP2) antigen in cellular

lysates of S-HVT-003 infected cells (42kD present) and S-HVT-001 infected cells (42kD negative). IBDV specific polypeptides were identified using a VP2 specific rabit anti-peptide antiserum. The lanes contain: 1) protein molecular weight standards, 2) wild-type HVT infected CEF's, 3) S-HVT-001 infected CEF's, 4) S-HVT-003 infected CEF's, 5) S-HVT-003 infected CEF's, and 6) IBDV virion polypeptides.

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## Figures 6A-6C: Details of S-HVT-004 Construction.

Figure 6A is a restriction map of HVT DNA in the region of the BamHI #16 fragment. This fragment is contained within a large HindIII fragment. Shown also is the XhoI site (X) where applicants have made their insertion. Before the insertion, the XhoI was first changed to EcoRI (R) site by use of a "linker" and standard cloning procedures.

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Figure 6B provides details of the construction of the  $\beta$ -gal gene and MDV gA gene inserted into the BamHI #16 fragment for use in homologous recombination. Beta-gal was under the control of the PRV gX gene promoter (gX), while the MDV gA gene was under the control of its own promoter.

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Figure 6C is of S-HVT-004 genome showing the location of the two inserted foreign genes,  $\beta$ -gal and MDV gA.

In Figure 6, H = HindIII; B = BamHI; X = XhoI; R
= EcoRI; Xb = XbaI; UL = unique long region; US =
unique short region; IR = internal repeat region;
TR = terminal repeat region.

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#### Figures 7A-7B:

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Detailed description of the  $\beta$ -galactosidase (lacZ) marker gene insertion in homology vector 467-22.A12. Figure 7A shows a diagram indicating the orientation of DNA fragments assembled in the marker gene. The origin of each fragment is described in the Materials and Methods section. Figures 7A and 7B show the DNA sequences located at the junctions between DNA fragments and at the ends of the marker gene (SEQ ID NOs: 28, 29, 30, 31, 32, and 33). Figures 7A and 7B further show the restriction sites used to generate each DNA fragment at the appropriate junction and the location of the lacZ gene coding region. Numbers in parenthesis () refer to amino acids, restriction sites in brackets [] indicate the remnants of sites which were destroyed during construction. The following abbreviations are used, pseudorabies virus (PRV), lactose operon Z gene (lacZ), Escherichia coli (E.Coli). polyadenylation signal (pA), and glycoprotein X (gpX).

#### Figure 8:

BamHI, NotI restriction map of the HVT genome. 25 The unique long (UL) and unique short (US) regions are shown. The long and short region repeats are indicated by boxes. The BamHI fragments are numbered in decreasing order of size. The 30 location of probes P1-P4 are indicated. The origin of each probe is as follows: P1 - BamHI #6, P2 - BamHI #2, P3 - BamHI #13, and P4 - 4.0 kb BgIII to StuI sub-fragment of HVT genomic XbaI fragment #5 (8.0 kb).

Figure 9: Shows the Procedure for construction of plasmid pSY229.

#### Figures 10A-10B:

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Detailed description of the MDV gene cassette insert in Homology Vectors 456-18.18 and 456-Figure 10A and 10B show a diagram indicating the orientation of DNA fragments assembled in the cassette and the location of the MDV gA and gB genes. The origin of each fragment is described in the Materials and Methods section. The sequences located at the junctions between each fragment and at the ends of the marker gene are shown in Figures 10A and 10B, including junction A (SEQ ID NO: 34), junction B (SEQ ID NO: 35), and junction C (SEQ ID NO: 36). restriction sites used to generate each fragment indicated at are the appropriate junction. Numbers in parenthesis () refer to amino acids, and restriction sites in brackets [] indicate the remnants of sites which were destroyed during construction.

#### Figures 11A-11B:

Detailed description of the HindIII fragment insert in Homology Vector 556-41.5. The diagram of Figures 11A and 11B show the orientation of DNA fragments assembled in the cassette. The origin of each fragment is described in the Materials and Methods section. Figures 11A and 11B further show the DNA sequences located at the junctions between each DNA fragment of the plasmid and at the ends of the marker gene, including junction A (SEQ ID NO: 37), junction B (SEQ ID NO: 38), and junction C (SEQ ID NO: 39). The restriction sites used to generate each fragment are indicated at the appropriate junction. The location of the MDV gD

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and a portion of the gI gene\_is also given. Numbers in parenthesis () refer to amino acids, and restriction sites in brackets [] indicate the remnants of sites which were destroyed during construction.

#### Figures 12A-12C:

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Detailed description of the SalI fragment insert in Homology Vector 255-18.B16. Figure 12A shows diagram indicating the orientation of DNA fragments assembled in the cassette. The origin of each fragment is described in the Materials and Methods section. Figures 12A to 12C further show the DNA sequences located at the junctions between each fragment and at the ends of the marker gene are shown, including junction A (SEQ ID NO: 40), junction B (SEQ ID NO: 41), junction C (SEQ ID NO: 42), junction D (SEQ ID NO: 43), junction E (SEQ ID NO: 44), junction F(SEQ ID NO: 45), junction G (SEQ ID NO: 46), and junction H (SEQ ID NO: 47). restriction sites used to generate each fragment are indicated at the appropriate junction. The location of the NDV F and lacZ-NDV HN hybrid gene are shown. Numbers in parenthesis () refer to amino acids, and restriction sites in brackets [] indicate the remnants of sites which were destroyed during construction.

#### Figures 13A-13B:

Show how the unique XhoI site of the BamHI #10 fragment of the HVT genome was converted into a PacI site and a NotI site by insertion of the synthetic DNA sequence at the XhoI site (Nucleotides #1333-1338; SEQ ID NO. 48). Figure 13A shows the Xho site converted into a PacI site to generate Plasmid 654-45.1 (SEQ ID NO. 55) and Figure 13B shows the XhoI site converted into a

NotI site to generate Plamid 686-63.A1 (SEQ ID NO. 56).

#### Figure 14:

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Restriction map and open reading frames of the sequence surrounding the insertion site within the unique long of HVT (SEQ ID NO. 48). This map shows the XhoI restriction site (SEQ ID NO. 48; Nucl. 1333-1338) used for insertion of foreign genes. Also shown are four open reading frames within this sequence. ORF A is interrupted by insertion of DNA into the XhoI site. The ORF A amino acid sequence (SEQ ID NO. 50; Nucl. 1402 to 602; 267 acids) shows no significant sequence identity to any known amino acid sequence in the protein databases. UL 54 (SEQ ID NO. 49; Nucl. 146 to 481; 112 amino acids) and UL55 (SEQ ID NO. 51; Nucl. 1599 to 2135; 179 amino acids) significant sequence identity to the herpes simplex virus type I UL54 and UL55 proteins, respectively. ORF B (SEQ ID NO. 52; Nucl. 2634 to 2308; 109 amino acids) shows no significant sequence identity to any known amino acid sequence in the protein databases. Searches were performed on NCBI databases using Blast software.

#### Figure 15:

Restriction map of cosmids 407-32.1C1, 672-01.A40, 672-07.C40, and 654-45.1. The overlap of HVT genomic DNA fragments EcoRI #9 and BamHI #10 is illustrated. A unique XhoI site within the EcoRI #9 and BamHI #10 fragments has been converted to a unique PacI site in Plasmid 654-45.1 or a unique NotI site in Plasmid 686-63.A1.

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#### DETAILED DESCRIPTION OF THE INVENTION

This invention provides a recombinant herpesvirus of turkeys (HVT) comprising a foreign DNA sequence inserted into a non-essential site in the HVT genome. The foreign DNA sequence is capable of being expressed in a host cell infected with the recombinant HVT and its expression is under the control of a promoter located upstream of the foreign DNA sequence.

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As defined herein "a non-essential site in the HVT genome" means a region in the HVT viral genome which is not necessary for the viral infection or replication.

As defined herein, "viral genome" or "genomic DNA" means the entire DNA which the naturally occurring herpesvirus of turkeys contains. As defined herein, "foreign DNA sequence" or "gene" means any DNA or gene that is exogenous to the genomic DNA.

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As defined herein, an "open reading frame" is a segment of DNA which contains codons that can be transcribed into RNA which can be translated into an amino acid sequence and which does not contain a termination codon.

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The invention further provides several appropriate insertion sites in the HVT genome useful for constructing the recombinant herpesvirus of the present invention. Insertion sites include the EcoRI #9 fragment and the BamHI #10 fragment of the HVT genome, a preferred insertion site within both of those fragments being a XhoI restriction endonuclease.

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Another such site is the BamHI #16 fragment of the HVT genome. A preferred insertion site within the BamHI #16 fragment lies within an open reading frame encoding

UL43 protein and a preferred insertion site within that open reading frame in a *XhoI* restriction endonuclease site.

Yet another insertion site is the HVT US2 gene, with a preferred insertion site within it being a StuI endonuclease site.

This invention provides a recombinant herpesvirus of turkeys comprising a herpesvirus of turkeys viral genome which contains a foreign DNA sequence inserted within the EcoRl #9 fragment of the herpesvirus of turkeys viral genome, and the foreign DNA sequence is capable of being expressed in a host cell infected with the herpesvirus of turkeys.

In one embodiment, the foreign DNA sequence is inserted within an Open Reading Frame A (ORFA) of the EcoR1 #9 fragment. Insertion of foreign DNA sequences into the XhoI site of EcoR1 #9 interrupts ORFA indicated that the entire ORFA region is non-essential for replication of the recombinant.

purposes of this invention, "a recombinant herpesvirus of turkeys" is a live herpesvirus of turkeys which has been generated by the recombinant methods well known to those of skill in the art, e.g., methods set forth in DNA TRANSFECTION GENERATING RECOMBINANT HERPESVIRUS in Materials and Methods, and the virus has not had genetic material essential for the replication of the herpesvirus of turkeys deleted. The purified herpesvirus of turkeys results in stable insertion of foreign DNA sequences or a gene in the EcoR1 #9 fragment or BamH1 #10 fragment.

The invention further provides recombinant herpesvirus of turkeys where the foreign DNA sequence encodes a

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polypeptide which is antigenic in an animal into which the recombinant herpesvirus is introduced.

In one embodiment the polypeptide is a detectable marker. For purposes of this invention, a "polypeptide 5 which is a detectable marker" includes the bimer, trimer and tetramer form of the polypeptide. E. coli B-galactosidase is а tetramer composed polypeptides or monomer subunits. In one embodiment 10 polypeptide coli beta-galactosidase. is E . Preferably this recombinant herpesvirus of turkeys is designated S-HVT-001, S-HVT-014, or S-HVT-012.

S-HVT-012 has been deposited on October 15, 1992

pursuant to the Budapest Treaty on the International Deposit of Microorganism for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2382.

S-HVT-014 has been deposited on December 7, 1993 pursuant to the Budapest Treaty on the International Deposit of Microorganism for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2440.

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In another embodiment the foreign DNA sequence encodes a cytokine. In another embodiment the cytokine is chicken myelomonocytic growth factor (cMGF) or chicken interferon (cIFN). In a preferred embodiment the recombinant herpesvirus of turkeys is designated S-HVT
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The invention further provides a recombinant

herpesvirus of turkeys whose viral genome contains foreign DNA encoding an antigenic polypeptide which is from Marek's disease virus (MDV), Newcastle disease virus (NDV), infectious laryngotracheitis virus (ILTV), infectious bronchitis virus (IBV) or infectious bursal disease virus (IBDV).

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This invention provides a recombinant herpesvirus of turkeys with a foreign DNA sequence insertion in the EcoR1 #9 fragment which further comprises a foreign DNA sequence encoding the antigenic polypeptide selected from the group consisting of: Marek's disease virus, Newcastle disease virus, infectious laryngotracheitis virus, infectious bronchitis virus and infectious bursal disease virus.

In one embodiment the foreign DNA sequence encoding the antigenic polypeptide is from Marek's disease virus and encodes Marek's disease virus glycoprotein gA, Marek's disease virus glycoprotein gB or Marek's disease virus glycoprotein gD. In another embodiment the foreign DNA sequences encoding the Marek's disease virus glycoprotein gA, glycoprotein gB or glycoprotein gD are inserted into the unique StuI site of the US2 gene coding region of the herpesvirus of turkeys.

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding antigenic polypeptide from Marek's disease virus. Preferably, the antigenic polypeptide is Marek's disease virus glycoprotein gB, gA or gD.

In one embodiment a recombinant HVT containing a foreign DNA sequence encodes IBDV VP2, MDV gA, and MDV gB. Preferably, such recombinant virus is designated S-HVT-137 and S-HVT-143.

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The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gA and further comprising foreign DNA encoding a polypeptide which is a detectable marker. Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-004.

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gB. Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-045.

An embodiment of a recombinant HVT containing a foreign DNA sequence encoding MDV gB is also provided and this recombinant HVT is designated S-HVT-045. S-HVT-045 has been deposited on October 15, 1992 pursuant to the Budapest Treaty on the International Deposit of Microorganism for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2383.

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The present invention also provides recombinant HVTs engineered to contain more than one foreign DNA sequence encoding an MDV antigen. For example, a foreign DNA sequence encoding MDV gA and gB can both be vectored into the HVT genome. Furthermore, a recombinant HVT can be constructed to include a foreign DNA sequence encoding MDV gA, gB, and gD.

Recombinant HVT designated S-HVT-046 and S-HVT-047
provide embodiments of a recombinant HVT containing foreign DNA sequence encoding MDV gA and gB; recombinant HVT designated S-HVT-048 and S-HVT-062

provide embodiments of a recombinant HVT containing foreign DNA sequence encoding MDV qA, qB and qD.

S-HVT-062 has been deposited on February 23, 1993 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Paten Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2401.

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The present invention provides a recombinant HVT containing a foreign DNA sequence encoding an antigenic polypeptide from Newcastle disease virus (NDV). In such case, it is preferred that the antigenic polypeptide is Newcastle disease virus fusion (F) protein or Newcastle disease virus hemagglutininneuraminidase (HN), or a recombinant protein comprising E. coli B-galactosidase fused to Newcastle disease virus hemagglutinin-neuraminidase (HN). One example of such a virus is designated S-HVT-007.

The present invention also provides recombinant HVTs engineered to contain one or more foreign DNA sequence encoding an antigenic polypeptide form MDV as well as one or more foreign DNA sequences encoding an antigenic polypeptide from NDV. Preferably, the MDV antigenic polypeptide is MDV gB, gD, or gA and the NDV F or HN.

In one embodiment of the invention, the recombinant HVT contains foreign DNA sequence encoding MDV gB, MDV gA and NDV F. Preferably, this HVT is designated S-HVT-048.

In one embodiment of the invention, the recombinant HVT contains foreign DNA sequence encoding MDV gB, MDV gA and NDV HN. Preferably, this HVT is designated S-HVT-

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For example, a foreign DNA sequence encoding MDV gA and gB can both be vectored into the HVT genome. Furthermore, a recombinant HVT can be constructed to include a foreign DNA sequence encoding MDV gA, gB, and gD.

Further, in another embodiment the foreign DNA sequence encoding the antigenic polypeptide is from Newcastle 10 disease virus and encodes Newcastle disease virus fusion protein or Newcastle disease virus hemagglutinin-neuraminidase. In another embodiment the foreign DNA sequences encoding the Newcastle disease 15 virus fusion protein or Newcastle disease virus hemagglutinin-neuraminidase are inserted into a XhoI site in EcoR1 #9 of the unique long region of the herpesvirus of turkeys. In a preferred embodiment the recombinant herpesvirus of turkeys is designated S-HVT-20 136.

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding antigenic polypeptide from Marek's disease virus and further comprising foreign DNA encoding antigenic polypeptide form Newcastle disease virus.

The present invention further provides a recombinant HVT which contains a foreign DNA sequence encoding an antigenic polypeptide from Marek's disease virus glycoprotein gB and Marek's disease virus glycoprotein and further comprising foreign DNA encoding Newcastle disease virus fusion (F) protein. Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-048.

The invention further provides recombinant herpesvirus

of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gB and Marek's disease virus glycoprotein gA and further comprising foreign DNA encoding Newcastle disease virus hemagglutinin-neuraminidase (HN). Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-049.

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gB and Marek's disease virus glycoprotein gA and further comprising foreign DNa encoding Newcastle disease virus fusion (F) protein and Newcastle disease virus hemagglutinin-neuraminidase (HN). Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-050.

S-HVT-050 has been deposited on February 23, 1993 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purpose of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2400.

In yet another embodiment of the invention, the recombinant HVT contains foreign DNA sequence encoding MDV gB, MDV gA, MDV gD, NDV F and NDV HN. Preferably, such recombinant HVT is designated S-HVT-106 or S-HVT 128.

The invention further provides recombinant herpesvirus Further, in one embodiment the foreign DNA sequence encodes the antigenic polypeptide from an infectious laryngotracheitis virus and encodes infectious laryngotracheitis virus glycoprotein gB, infectious

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laryngotracheitis virus glycoprotein gI or infectious laryngotracheitis virus glycoprotein gD.

In another embodiment the foreign DNA sequence encodes an antigenic polypeptide which is derived or derivable from a group consisting of: MDV gA, MDV gB, MDV gD, NDV HN, NDV F, ILT gB, ILT gI, ILT gD, IBV, IBDV VP2, IBDV VP3, IBDV VP4, avian encephalomyelitis virus, avian reovirus, avian paramyxovirus, avian influenza virus, avian adenovirus, fowl pox virus, avian coronavirus, avian rotavirus, chick anemia virus (agent), Salmonella spp. E. coli, Pasteurella spp., Bordetella spp., Eimeria spp., Histomonas spp., Trichomonas spp., Poultry nematodes, cestodes, trematodes, poultry mites/lice, poultry protozoa.

The invention further provides a recombinant herpesvirus of turkeys which contains a foreign DNA sequence encoding an antigenic polypeptide from infectious laryngotracheitis virus. It is preferred that the antigenic polypeptide is ILTV glycoprotein gB, ILTV gD or ILTV gI.

Also provided are recombinant HVTs which are engineered to contain more than one foreign DNA sequence encoding an ILTV antigen. For example, ILTV gB and gD can be vectored together into the HVT genome, so can ILTV gD and gI, and ILTV gB, gD and gI. Recombinant HVT designated S-HVT-051, S-HVT-052, and S-HVT-138 are embodiments of such recombinant virus.

The present invention also provides a recombinant HVT which contains more than one foreign DNA sequence encoding an antigenic polypeptide from MDV as well as one or more foreign DNA sequences encoding an antigenic polypeptide from ILTV. Preferably, the MDV antigenic polypeptide is MDV gB, gD or gA and the ILTV antigenic

polypeptide is ILTV gB, gD or gI.

In one embodiment of the invention, the recombinant HVT contains foreign DNA sequences encoding MDV gB, MDV gA, MDV gD, ILTV gD and ILTV gB. Preferably, this recombinant HVT is designated S-HVT-123.

In another embodiment of this invention, the recombinant HVT contains foreign DNA sequences encoding MDV gB, MDV gA, MDV gD, ILTV gland ILTV gD. Preferably, this recombinant HVT is designated S-HVT-139 or S-HVT-140.

of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gB, Mareck's disease virus glycoprotein gA, and Marek's disease virus glycoprotein gD and further comprising foreign DNA which encodes infectious laryngotracheitis virus glycoprotein gD, infectious laryngotracheitis virus glycoprotein gB, and E. coli B-galactosidase. Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-104.

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding infectious bronchitis virus spike protein or infectious bronchitis virus matrix protein.

The present invention further provides a recombinant HVT which contains a foreign DNA sequence encoding an antigenic polypeptide from infectious bronchitis virus (IBV). Preferably, the antigenic polypeptide is IBV spike protein or IBV matrix protein.

The present invention also provides a recombinant HVT which contains one or more foreign DNA sequences

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encoding an antigenic polypeptide from IBV as well as one or more foreign DNA sequences encoding an antigenic polypeptide from MDV. Preferably, the IBV antigenic polypeptide is IBV spike protein or IBV matrix protein, and the MDV antigenic polypeptide is MDV gB, gD or gA. One embodiment of such recombinant virus is designated S-HVT-066.

The invention further provides a recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding antigenic polypeptide from infectious bursal disease virus and further comprising foreign DNA encoding a polypeptide which is a detectable marker.

Further, in one embodiment a foreign DNA sequence encoding the antigenic polypeptide is from infectious bursal disease virus. In another embodiment the foreign DNA sequence encodes infectious bursal disease virus VP2 gene. In another embodiment the foreign DNA sequence encodes infectious bursal disease virus VP3 gene. In another embodiment the foreign DNA sequence encodes infectious bursal disease virus VP4 gene. Preferably, this recombinant herpesvirus of turkeys is designated S-HVT-003 or S-HVT-096.

Recombinant HVT designated S-HVT-003 and S-HVT-096 are each an embodiment of a recombinant HVT comprising foreign DNA sequence encoding antigenic polypeptide from IBDV and encoding a detectable marker. S-HVT-003 has been deposited on July 21, 1987 pursuant to the Budapest Treaty on the International Deposit of Microorganism for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR. 2178.

This invention provides a recombinant herpesvirus of turkeys containing a foreign DNA sequence inserted into the EcoR1 #9 fragment herpesvirus of turkeys viral genome wherein the foreign DNA sequence is from an infectious laryngotracheitis virus and encodes infectious laryngotracheitis virus glycoprotein gB, or infectious laryngotracheitis virus glycoprotein gD.

In one embodiment the foreign DNA sequence is from an infectious laryngotracheitis virus and encodes infectious laryngotracheitis virus glycoprotein gD, or laryngotracheitis virus glycoprotein gI.

This invention provides a recombinant herpesvirus of turkeys containing a foreign DNA sequence inserted into the EcoR1 #9 fragment herpesvirus of turkeys viral genome wherein the foreign DNA sequence is from an Newcastle disease virus and encodes a Newcastle disease virus HN or Newcastle disease virus F.

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This invention provides a recombinant herpesvirus of turkeys containing a foreign DNA sequence inserted into the EcoR1 #9 fragment herpesvirus of turkeys viral genome wherein the foreign DNA sequence is from an infectious bursal virus and encodes an infectious bursal disease virus VP2, VP3, VP4.

This invention provides a recombinant herpesvirus of turkeys containing a foreign DNA sequence inserted into the EcoR1 #9 fragment herpesvirus of turkeys viral genome wherein the foreign DNA sequence is from an infectious bronchitis virus and encodes an infectious bronchitis virus matrix protien.

In another embodiment a foreign DNA sequence encodes an antigenic polypeptide which is derived or derivable from a group consisting of: MDV gA, MDV gB, MDV gD, NDV

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HN, NDV F, ILT gB, ILT gI, ILT gD, IBV, IBDV VP2, IBDV VPD3, IBDV VP4, avian encephalomyelitis virus, avian reovirus, avian paramyxovirus, avian influenza virus, avian adenovirus, fowl pox virus, avian coronavirus. avian rotavirus, chick anemia virus (agent), Salmonella coli, Pasteurella spp., Bordetella spp., spp. E. Eimeria spp., Histomonas spp., Trichomonas spp., Poultry cestodes, trematodes, nematodes, mites/lice, poultry protozoa. In a preferred embodiment the recombinant herpesvirus of turkeys is designated S-HVT-136.

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Such antigenic polypeptide may be derived or derivable from the following: feline pathogen, canine pathogen, equine pathogen, bovine pathogen, avian pathogen, porcine pathogen, or human pathogen.

In another embodiment, the antigenic polypeptide of a human pathogen is derived from human herpesvirus, herpes simplex virus-1, herpes simplex virus-2, human cytomegalovirus, Epstein-Barr virus, Varicell-Zoster virus, human herpesvirus-6, human herpesvirus-7, human influenza, human immunodeficiency virus, rabies virus, measles virus, hepatitis B virus and hepatitis C virus. Furthermore, the antigenic polypeptide of a human pathogen may be associated with malaria or malignant tumor from the group consisting of Plasmodium falciparum, Bordetella pertusis, and malignant tumor.

The invention further provides recombinant herpes virus of turkeys whose genomic DNA contains foreign DNA encoding Newcastle disease virus fusion (F) protein and further comprising foreign DNA encoding a recombinant protein, wherein E. coli B-galactosidase is fused to Newcastle disease virus hemagglutinin-neuraminidase (HN).

The invention further provides recombinant herpesvirus of turkeys whose genomic DNA contains foreign DNA encoding Marek's disease virus glycoprotein gB and Marek's disease virus glycoprotein gA and further comprising foreign DNA encoding Newcastle disease virus hemagglutinin-neuraminidase (HN).

This invention provides a recombinant herpesvirus of turkeys-Marek's disease virus chimera comprising a herpesvirus of turkeys unique long viral genome region and a Marek's disease virus unique short region. In one embodiment the recombinant herpesvirus of turkeys-Marek's disease virus chimera contains a foreign DNA sequence inserted within the EcoR1 #9 fragment of the herpesvirus of turkeys viral genome, and the foreign DNA sequence capable of being expressed in a host cell'infected with the herpesvirus of turkeys.

In one embodiment the recombinant herpesvirus of turkeys contains a foreign DNA sequence which encodes a polypeptide. The polypeptide may be antigenic in an animal into which the recombinant herpesvirus is introduced.

In another embodiment the polypeptide is *E. coli* betagalactosidase. In another embodiment the foreign DNA sequence encodes a cytokine. In another embodiment the cytokine is chicken mylomonocytic growth factor (cMGF) or chicken interferon (cIFN).

The invention further provides recombinant herpesvirus of turkeys where the foreign DNA sequence encodes a polypeptide which is antigenic in an animal into which the recombinant herpesvirus is introduced.

Further, the recombinant herpesvirus of turkeys further comprises a foreign DNA sequence encoding the antigenic

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polypeptide selected from the group consisting of: Marek's disease virus, Newcastle disease virus, infectious laryngotracheitis virus, infectious bronchitis virus and infectious bursal disease virus.

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This invention provides a recombinant herpesvirus of turkeys wherein the foreign DNA sequence is under control of an endogenous upstream herpesvirus promoter. In one embodiment the foreign DNA sequence is under control of a heterologous upstream promoter. In another embodiment the promoter is selected from PRV gX, HSV-1 alpha 4, HCMV immediate early, MDV gA, MDV gB, MDV gD, ILT gB, BHV-1.1 VP8 and ILT gD.

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This invention provides a homology vector for producing a recombinant herpesvirus of turkeys by inserting foreign DNA into the viral genome of a herpesvirus of turkey which comprises a double-stranded DNA molecule consisting essentially of: a) double stranded foreign DNA not usually present within the herpesvirus of turkeys viral genome; b) at one end the foreign DNA, double-stranded herpesvirus of turkeys DNA homologous to the viral genome located at one side of the EcoR1 #9 site the coding region of the herpesvirus of turkeys viral genome; and c) at the other end of the foreign double-stranded herpesvirus of turkeys DNA. homologous to the viral genome located at the other side of the EcoR1 #9 fragment of the coding region of the herpesvirus of turkeys viral genome. Examples of the homology vectors are designated 751-87.A8 and 761-7.A1.

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In one embodiment the polypeptide is antigenic in the animal into which the recombinant herpesvirus of turkeys is introduced. In another embodiment the antigenic polypeptide is from a cytokine, Marek's disease virus, Newcastle disease virus, infectious

laryngotracheitis virus, or infectious bronchitis a preferred embodiment the antiqenic In virus. polypeptide is a chicken mylomonocytic growth factor (cMGF) or chicken interferon (cIFN), infectious bursal disease virus polyprotein, infectious bursal disease virus VP2 protein, Marek's disease virus glycoprotein qB, Marek's disease virus glycoprotein gA, Marek's disease virus glycoprotein gD, Newcastle disease virus fusion protein, Newcastle disease virus hemagglutininlaryngotracheitis neuraminidase, infectious glycoprotein gB, infectious laryngotracheitis virus glycoprotein gD, infectious bronchitis virus spike protein, or infectious bronchitis virus matrix protein.

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15 In another embodiment the double stranded foreign DNA sequence in the homology vector encodes an antigenic polypeptide derived from an equine pathogen. The antiquenic polypeptide of an equine pathogen can derived from equine influenza virus or equine herpesvirus. Examples of such antigenic polypeptide are equine 20 influenza virus type A/Alaska 91 neuraminidase, equine influenza virus type A/Prague 56 neuraminidase, equine influenza virus type A/Miami 63 neuraminidase, equine influenza virus type A/Kentucky 81 neuraminidaseequine 25 glycoprotein herpesvirus type 1 B, and equine herpesvirus type 1 glycoprotein D.

> In another embodiment the double stranded foreign DNA sequence of the homology vector encodes an antigenic polypeptide derived from bovine respiratory syncytial virus or bovine parainfluenza virus. The antigenic polypeptide of derived from bovine respiratory syncytial virus equine pathogen can derived from equine influenza virus is bovine respiratory syncytial virus (BRSV G), bovine attachment protein respiratory syncytial virus fusion protein (BRSV F), respiratory syncytial virus nucleocapsid protein (BRSV

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N), bovine parainfluenza virus type 3 fusion protein, and the bovine parainfluenza virus type 3 hemagglutinin neuraminidase.

In another embodiment the double stranded foreign DNA sequence in the homology vector encodes a cytokine capable of stimulating human immune response. For example, the cytokine may be, but is not limited to, interleukin-2, interleukin-6, interleukin-12, interferons, granulocyte-macrophage colony stimulating factors, and interleukin receptors.

In one embodiment of the invention, the double-stranded herpesvirus of turkeys DNA is homologous to DNA sequences present within the BamHI #16 fragment of the herpesvirus of turkeys genome. Preferably, the double-stranded herpesvirus of turkeys DNA is homologous to DNA sequences present within the open reading frame encoding UL 43 protein of the herpesvirus of turkeys genome. Preferably, this homology vector is designated 172-29.31.

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For purposes of this invention, a "homology vector" is a plasmid constructed to insert foreign DNA in a specific site on the genome of a herpesvirus of turkeys.

In one embodiment of the invention, the double-stranded herpesvirus of turkeys DNA is homologous to DNA sequences present within the *EcoR1* #9 fragment of the herpesvirus of turkeys genome. Preferably, this homology vector is designated 172-63.1.

In one embodiment of the invention, the double-stranded herpesvirus of turkeys DNA is homologous to DNA sequences present within the US2 gene coding region of the herpesvirus of turkeys genome. Preferably, this homology vector is designated 435-47.1.

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In another embodiment the foreign DNA sequence encodes a screenable marker. Examples of screenable markers, inlcude but are not limited to: E. coli B-galactosidase or E. coli B-glucuronidase.

The invention further provides a vaccine which comprises an effective immunizing amount of a recombinant herpesvirus of turkeys of the present invention and a suitable carrier.

This invention provides a vaccine useful for immunizing a bird against Marek's disease virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a vaccine useful for immunizing a bird against Newcastle disease virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a vaccine useful for immunizing a bird against infectious laryngotracheitis virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a vaccine useful for immunizing a bird against infectious bronchitis virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a vaccine useful for immunizing a bird against infectious bursal disease virus which comprises an effective immunizing amount of the

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recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a multivalent vaccine useful for immunizing a bird against Marek's disease virus and Newcastle disease virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys.

This invention provides a multivalent vaccine useful for immunizing a bird against Marek's disease virus and infectious laryngotracheitis virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

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This invention provides a multivalent vaccine useful for immunizing a bird against Marek's disease virus and infectious bronchitis virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

This invention provides a multivalent vaccine useful for immunizing a bird against Marek's disease virus and infectious bursal disease virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys and a suitable carrier.

The present invention also provides a method of immunizing a fowl. For purposes of this invention, 30 this includes immunizing a fowl against infectious bursal disease virus, Marek's disease virus, Newcastle disease virus, infectious laryngotracheitis virus, or infectious bronchitis virus. The method comprises administering to the fowl an effective immunizing dose 35 of the vaccine of the present invention. The vaccine may be administered by any of the methods well known to those skilled in the art. for example. by

intramuscular, subcutaneous, intraperitoneal or intravenous injection. Alternatively, the vaccine may be administered intranasally or orally.

5 This invention provides a host cell infected with the recombinant herpesvirus of turkey. In one embodiment the host cell is an avian cell.

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For purposes of this invention, a "host cell" is a cell used to propagate a vector and its insert. Infecting the cell was accomplished by methods well known to those skilled in the art, for example, as set forth in DNA TRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUS in Materials and Methods. Methods for constructing, selecting and purifying recombinant herpesvirus of turkeys are detailed below in

This invention provides a method of distinguishing chickens or other poultry which are vaccinated with the above vaccine from those which are infected with a naturally-occurring Marek's disease virus comprises analyzing samples of body fluids from chickens or other poultry for the presence glycoprotein gG and at least one other antigen normally expressed in chickens or other poultry infected by a naturally-occurring Marek's disease virus, the presence of those antigens normally expressed in infected chickens but the absence of glycoprotein qG being indicative of vaccination with the above vaccine and not infection with a naturally-occurring Marek's disease virus.

This invention provides a recombinant herpesvirus of turkeys which expresses foreign DNA sequences is useful as vaccines in avian or mammalian species including but not limited to chickens, turkeys, ducks, feline, canine, bovine, equine, and primate, including human.

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This vaccine may contain either inactivated or live recombinant virus.

purposes of this invention, an "effective immunizing amount" of the recombinant feline herpes virus of the present invention is within the range of 10° PFU/dose. In another embodiment immunizing amount is  $10^5$  to  $10^7$  PFU/dose. In a preferred embodiment the immunizing amount is  $10^{6}$ PFU/dose.

The method comprises administering to the animal an effective immunizing dose of the vaccine of the present The vaccine may be administered by any of invention. the methods well known to those skilled in the art, for example, by intramuscular, subcutaneous, intraperitoneal or intravenous injection. Alternatively, the vaccine may administered be intranasally or orally.

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Suitable carriers for the recombinant virus are well known to those skilled in the art and include but are not limited to proteins, sugars, etc. One example of such a suitable carrier is a physiologically balanced culture medium containing one or more stabilizing agents such as hydrolyzed proteins, lactose, etc. Preferably, the live vaccine is created by taking tissue culture fluids and adding stabilizing agents such as stabilizing, hydrolyzed proteins. Preferably, the inactivated vaccine uses tissue culture fluids directly after inactivation of the virus.

This invention is further illustrated in the Experimental Details section which follows. This section is set forth to aid in an understanding of the invention but is not intended to, and should not be construed to, limit in any way the invention as set

forth in the claims which follow thereafter.

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## EXPERIMENTAL DETAILS:

## Materials and Methods

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PREPARATION OF HERPESVIRUS OF TURKEYS STOCK SAMPLES. Herpesvirus of turkeys stock samples were prepared by infecting tissue culture cells at a multiplicity of infection of 0.01 PFU/cell in Dulbecco's Modified Eagle Medium (DMEM) containing 2 mM glutamine, 100 units/ml penicillin, 100 units/ml streptomycin (these components are obtained from Irvine Scientific or an equivalent supplier, and hereafter are referred to as complete DME medium) plus 1% fetal bovine serum. After cytopathic effect was complete, the medium and cells were harvested and the cells were pelleted at 3000 rpm for 5 minutes in a clinical centrifuge. Infected cells were resuspended in complete medium containing 20% fetal bovine serum, 10% DMSO and stored frozen at -70°C.

PREPARATION OF HERPESVIRUS OF TURKEY DNA. All manipulations of herpesvirus of turkey (HVT) were made using strain FC-126 (ATCC #584-C). For the preparation of HVT viral DNA from the cytoplasm of infected cells, primary chicken embryo fibroblasts were infected at a MOI sufficient to cause extensive cytopathic effect before the cells overgrew. All incubations were carried out at 39°C in a humidified incubator with 5% CO, in air. Best DNA yields were obtained by harvesting monolayers which were maximally infected, but showing incomplete cell lysis (typically 5-7 days). Infected cells were harvested by scraping the cells into the medium using a cell scraper (Costar brand). The cell suspension was centrifuged at 3000 rpm for 10 minutes at 5°C in a GS-3 rotor (Sorvall Instruments). The resultant pellet was resuspended in cold PBS (20

ml/Roller Bottle) and subjected to another centrifugation for 10 minutes at 3000 rpm in the cold. After decanting the PBS, the cellular pellet was resuspended in 4 ml/roller bottle of RSB buffer (10 mM Tris pH 7.5, 1 mM EDTA, and 1.5 mM MqCl<sub>2</sub>). (Nonidet P-40"; Sigma) was added to the sample to a final concentration of 0.5% minutes with occasional The sample was centrifuged for 10 minutes at 3000 rpm in the cold to pellet the nuclei and remove cellular debris. The supernatant fluid was carefully transferred to a 15 ml Corex centrifuge tube. EDTA (0.5M pH 8.0) and SDS (sodium dodecyl sulfate; stock 20%) were added to the sample to final concentrations of 5 mM and 1%, respectively. hundred  $\mu$ l of proteinase-K (10 mg/ml; Boehringer Mannheim) was added per 4 ml of sample, mixed, and incubated at 45°C for 1-2 hours. After this period, an equal volume of water-saturated phenol was added to the sample and gently mixed by hand. The sample was spun in a clinical centrifuge for 5 minutes at 3000 rpm to separate the phases. NaAc was added to the aqueous phase to a final concentration of 0.3M (stock solution 3M pH 5.2), and the nucleic acid precipitated at -70°C for 30 minutes after the addition of 2.5 volumes of cold absolute ethanol. DNA in the sample was pelleted by spinning for 20 minutes to 8000 rpm in an HB-4 rotor at 5°C. The supernatant was carefully removed and the DNA pellet washed once with 25 ml of 80% ethanol. DNA pellet was dried briefly by vacuum (2-3 minutes), and resuspended in 50  $\mu$ l/roller bottle of infected cells of TE buffer (10 mM Tris pH 7.5, 1 mM EDTA). Typically, yields of viral DNA ranged between 5-10 μg/roller bottle of infected cells. All viral DNA was stored at approximately 10°C.

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POLYMERASE FILL-IN REACTION. DNA was resuspended in buffer containing 50 mM Tris pH 7.4, 50 mM KCl, 5 mM

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MgCl<sub>2</sub>, and 400 micromolar each of the four deoxynucleotides. Ten units of Klenow DNA polymerase (BRL) were added and the reaction was allowed to proceed for 15 minutes at room temperature. The DNA was then phenol extracted and ethanol precipitated as above.

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DNA SEQUENCING. Sequencing was performed using the USB Sequenase Kit and 35S-dATP (NEN). Reactions using both the dGTP mixes and the dITP mixes were performed to clarify of areas compression. Alternatively. compressed areas were resolved on formamide gels. Templates were double-stranded plasmid subclones or single stranded M13 subclones, and primers were either made to the vector just outside the insert to be sequenced, or to previously obtained Sequence obtained was assembled and compared using Dnastar software. Manipulation and comparison of sequences obtained was performed with Superclone and Supersee programs from Coral Software.

MOLECULAR BIOLOGICAL TECHNIQUES. Techniques for the manipulation of bacteria and DNA, including such procedures as digestion with restriction endonucleases, gel electrophoresis, extraction of DNA from gels, ligation, phosphorylation with kinase, treatment with of phosphatase, growth bacterial cultures, transformation of bacteria with DNA, and molecular biological methods are described by Maniatis et al (1982) and Sambrook et al (1989). The polymerase chain reaction (PCR) was used to introduce restriction sites convenient for the manipulation of various DNAs. The procedures used are described by Innis et al (1990). In general amplified fragments were less than base pairs in size and critical regions of amplified fragments were confirmed by DNA sequencing. Except as noted, these techniques were used with minor

variation.

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SOUTHERN BLOTTING OF DNA. The general procedure for Southern blotting was taken from Maniatis et al. (1982). DNA was blotted to nitrocellulose filters (S&S BA85) in 20X SSC (1X ssc = 0.15M NaCl, 0.015M sodium citrate, pH 7.0), and prehybridized in hybridization solution consisting of 30% formamide, 1X Denhardt's solution (0.02% polyvinylpyrrolidone (PVP), bovine serum albumin (BSA), 0.02% Ficoll), 6X SSC, 50 mM NaH,PO, pH 6.8, 200  $\mu$ g/ml salmon sperm DNA for 4-24 hours at 55°C. Labeled probe DNA was added that had been labeled by nick translation using a kit from Bethesda Research Laboratories (BRL) and one 32P-labeled The probe DNA was separated from the nucleotide. unincorporated nucleotides by NACS column (BRL) or on a Sephadex G50 column (Pharmacia). After overnight hybridization at 55°C, the filter was washed once with 2X SSC at room temperature followed by two washes with 0.1% SSC, 0.1% sodium dodecyl sulfate (SDS) for 30 minutes at 55°C. The filter was dried and autoradiographed.

cDNA CLONING PROCEDURE. cDNA cloning refers to the methods used to convert RNA molecules into DNA molecules following state of the art procedures. Applicants' methods are described in (Gubler Hoffman. 1983). Bethesda Research Laboratories (Gaithersburg, MD) have designed a cDNA Cloning Kit that is very similar to the procedures used by applicants, and contains a set of reagents protocols that may be used to duplicate our results.

For cloning virus mRNA species, a host cell line sensitive to infection by the virus was infected at 5-10 plaque forming units per cell. When cytopathic effect was evident, but before total destruction, the

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medium was removed and the cells were lysed in 10 mls lysis buffer (4 M guanidine thiocyanate, 0.1% antifoam mM sodium citrate pH 7.0, 0.5% N-lauroyl sarcosine, 0.1 M beta-metcaptoethanol). lysate was poured into a sterilized Dounce homogenizer and homogenized on ice 8-10 times until the solution was homogenous. For RNA purification, 8 mls of cell lysate were gently layered over 3.5 mls of CsCl solution (5.7 M CsCl, 25 mM sodium citrate pH 7.0) in Beckman SW41 centrifuge tube. The samples were centrifuged for 18 hrs at 20° C at 36000 rpm in a Beckman SW41 rotor. The tubes were put on ice and the supernatants from the tubes were carefully removed by aspiration to leave the RNA pellet undisturbed. pellet was resuspended in 400 µl glass distilled water, and 2.6 mls of guanidine solution (7.5 M guanidine-HCL, 25 mM sodium citrate pH 7.0, 5 mM dithiothreitol) were The 0.37 volumes of 1 M acetic acid were added, followed by 0.75 volumes of cold ethanol and the sample was put at -20° C for 18 hrs to precipitate RNA. precipitate was collected by centrifugation in Sorvall centrifuge for 10 min a 4° C at 10000 rpm in an SS34 rotor. The pellet was dissolved in 1.0 ml distilled water, recentrifuged at 13000 rpm, and the supernatant saved. RNA was re-extracted from the pellet 2 more times as above with 0.5 ml distilled water, and the supernatants were pooled. A 0.1 volume of 2 M potassium acetate solution was added to the sample followed by 2 volumes of cold ethanol and the sample was put at -20° C for 18 hrs. The precipitated RNA was collected by centrifugation in the SS34 rotor at 4° C for 10 min at 10000 rpm. The pellet was dissolved in 1 ml distilled water and the concentration taken by absorption at A260/280. The RNA was stored at -70°C.

mRNA containing polyadenylate tails (poly-A) was

selected using oligo-dT cellulose (Pharmacia #27 5543-0). Three mg of total RNA was boiled and chilled and applied to the 100 mg oligo-dT cellulose column in binding buffer (0.1 M Tris pH 7.5, 0.5 M LiCl, 5mM EDTA pH 8.0, 0.1% lithium dodecyl sulfate). The retained poly-A RNA was eluted from the column with elution buffer (5mM Tris pH 7.5, 1mM EDTA pH 8.0, 0.1% sodium dodecyl sulfate). This mRNA was reapplied to an oligo-dT column in binding buffer and eluted again in elution buffer. The sample was precipitated with 200 mM sodium acetate and 2 volumes cold ethanol at -20°C for 18 hrs. The RNA was resuspended in 50  $\mu$ l distilled water.

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Ten  $\mu$ g poly-A RNA was denatured in 20 mM methyl mercury hydroxide for 6 min at 22°C. ß-mercaptoethanol was added to 75 mM and the sample was incubated for 5 min The reaction mixture for first strand cDNA synthesis in 0.25 ml contained 1  $\mu$ g oligo-dT primer (P-L Bio-chemicals) or 1  $\mu$ g synthetic primer, 28 units placental ribonuclease inhibitor (Bethesda Research Labs #5518SA), 100 mM Tris pH 8.3, 140 mM KCl, 10mM MgCl,, 0.8 mM dATP, dCTP, dGTP, and dTTP (Pharmacia), 100 microcuries 32p-labeled dCTP (New England Nuclear #NEG-013H), and 180 units AMV reverse transcriptase (Molecular Genetics Resources #MG 101). The reaction was incubated at 42°C for 90 min, and then was terminated with 20mM EDTA pH 8.0. The sample was extracted with an equal volume of phenol/chloroform (1:1) and precipitated with 2 M ammonium acetate and 2 volumes of cold ethanol -20°C for 3 hrs. precipitation and centrifugation, the pellet dissolved in 100  $\mu$ l distilled water. The sample was loaded onto a 15 ml G-100 Sephadex column (Pharmacia) in buffer (100 mM Tris pH 7.5, 1 mM EDTA pH 8.0, 100 mM The leading edge of the eluted DNA fractions was pooled, and DNA was concentrated by lyophilization until the volume was about 100  $\mu$ l, then the DNA was

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precipitated with ammonium acetate plus ethanol as above.

The entire first strand sample was used for second strand reaction which followed the Gubler and Hoffman (1983) method except that 50  $\mu$ g/ml dNTP's, 5.4 units DNA polymerase I (Boerhinger Mannheim #642-711), and 100 units/ml E. coli DNA ligase (New England Biolabs #205) in a total volume of 50 microliters were used. After second strand synthesis, the CDNA phenol/chloroform extracted and precipitated. was resuspended in 10  $\mu$ l distilled water, treated with 1 μg RNase A for 10 min at 22°C, and electrophoresed through a 1% agarose gel (Sigma Type II agarose) in 40 mM Tris-acetate pH 6.85. The gel was stained with ethidium bromide, and DNA in the expected size range was excised from the gel and electroeluted in 8 mM Tris-acetate Hq 6.85. Electroeluted DNA lyophilized to about 100 microliters, and precipitated with ammonium acetate and ethanol as above. was resuspended in 20  $\mu$ l water.

Oligo-dC tails were added to the DNA to facilitate The reaction contained the DNA, 100 mM potassium cacodylate pH 7.2, 0.2 mM dithiothreitol, 2mM and CaCl<sub>2</sub>, 80  $\mu$ moles dCTP, 25 units terminal transferase deoxynucleotidyl (Molecular Genetic Resources #S1001) in 50  $\mu$ l. After 30 min at 37°C, the reaction was terminated with 10mM EDTA, and the sample was phenol/chloroform extracted and precipitated as above.

The dC-tailed DNA sample was annealed to 200 ng plasmid vector pBR322 that contained oligo-dG tails (Bethesda Research Labs #5355 SA/SB) in 200  $\mu$ l of 0.01 M Tris pH 7.5, 0.1 M NaCl, 1 mM EDTA pH 8.0 at 65°C for 2 min and then 57°C for 2 hrs. Fresh competent E. coli DH-1

cells were prepared and transformed as described by Hanahan (1983) using half the annealed cDNA sample in twenty 200  $\mu$ l aliquots of cells. Transformed cells were plated on L-broth agar plates plus 10  $\mu$ g/ml tetracycline. Colonies were screened for the presence of inserts into the ampicillin gene using Ampscreen (Bethesda Research Labs #5537 UA), and the positive colonies were picked for analysis.

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FOR GENERATING RECOMBINANT 10 DNA TRANSFECTION HERPESVIRUS. The method is based upon the polybrene-DMSO procedure of Kawai and Nishizawa (1984) with the following modifications. Generation of recombinant HVT is dependent upon homologous recombination between HVT viral DNA and the plasmid homology vector 15 containing the desired foreign DNA flanked by the cloned herpesvirus sequences. appropriate Transfections were carried out in 6 cm plates (Corning plastic) of 50% confluent primary chick fibroblast (CEF) cells. The cells were plated out the 20 day before in CEF growth media (1X F10/199, 5% fetal calf serum, 2% glutamine, 1% non-essential amino acids, and 2% penicillin/streptomycin) containing 4 μg/ml mq/ml in lX polybrene (stock 4 HBSS). For cotransfections into CEF cells, 5 µg of intact HVT DNA, 25 and suspended in 1 ml of CEF media containing 30  $\mu$ g/ml polybrene (stock 4 mg/ml in 1X HBSS). polybrene suspension (1 ml) was then added to a 6 cm plate of CEF cells from which the media had been aspirated, and incubated at 39°C for 30 minutes. 30 The plates were rocked periodically during this time to redistribute the inoculum. After this period, 4 ml of CEF growth media was added directly to wash plate, and incubated an additional 2.5 hours a 39°C. time, the media was removed from each plate, and the 35 cells shocked with 2 ml of 30% DMSO Sulfoxide, J.T. Baker Chemical Co.) in 1X HBSS for 4

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minutes at room temperature. The 30% DMSO was carefully removed and the monolayers washed once with 1X HBSS at room temperature. The cells were then incubated at 39°C after the addition of 5 mls of CEF growth media. The next day, the media was changed to remove any last traces of DMSO and to stimulate cell Cytopathic effect from the virus becomes growth. apparent within 6 days. Generation of a high titer stock (80%-90% CPE) can usually be made within 1 week from this date. HVT stock samples were prepared by resuspending the infected cells in CEF growth media containing 20% fetal calf serum, 10% DMSO and stored at -70°C.

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15 PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The ability to generate herpesviruses by cotransfection of cloned overlapping subgenmoic fragments has been demonstrated pseudorabies virus (Zijl et al., 1988). If deletions 20 and/or insertions are engineered directly into the subgenomic fragments prior to the cotransfection, this procedure results in a high frequency of viruses containing the genomic alteration, greatly reducing the amount of screening required to purify the recombinant 25 virus. This procedure was used to construct recombinant HVT.

A library of subclones containing overlapping HVT subgenomic fragments was generated as follows. HVT DNA was obtained from the American Type Culture Collection (FC-126("Calnek")). It was sheared and then size selected on a glycerol gradient as described by van Zijl et al., (1988) with 40-50 kb fragments chosen as the insert population. The pooled fractions were diluted twofold with TE, one-tenth volume of 3M NaAc and 2.5 volumes of ethanol were added, and the DNA was precipitated at 30K rpm in a Beckman SW41 rotor for 1

The sheared fragments were given blunt ends by initial treatment with T4 DNA polymerase, using low DNTP concentrations to promote 3' overhang removal, followed by treatment with Klenow polymerase to fill in recessed 3' ends. These insert fragments were then ligated to a pWE15 (Strategene) cosmid vector, which had been digested with BamHI, treated with calf intestinal phosphatase, and made blunt by treatment with Klenow polymerase. The ligated mixture was then packaged using Gigapack XLpackaging extracts and (Stratagene). Ligation packaging was as recommended by the manufacturer.

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Published restriction maps for the enzymes BamHI, HindIII, and XhoI permitted the use of subcloned fragments as specific probes to screen the cosmid library for subclones spanning the genome. Probes were generated from subcloned restriction fragments. fragments were then labeled using a non-radioactive system (Genius, Boehringer Mannheim). Screening was facilitated by picking colonies to media followed by Sets of five filters and a master growth overnight. plate were stamped from microtiter dish and again grown overnight. Glycerol was added to the wells to 15% and the plates were frozen at -20°C to provide stock cultures of each colony. Filters were BioRad Colony Lift Membranes and were treated and hybridized per manufacturer's instructions, and washed in 0.1% SSC, 0.1% SDS, 65°C. Clones which hybridized with the nonradioactive probe were detected according to the Genius kit directions.

Colonies were selected for further analysis on the basis of their hybridization to two or more of the specific probes. These were then digested with BamHI, and compared to published maps of HVT (Buckmaster et al., 1988). The three cosmids (407-32.2C3,407-32.IG7,

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and 407-32.5G6) were obtained in this manner. A detailed description of each clone is given below. It was found that chloramphenicol amplification (Maniatis et al.,1982) was necessary to achieve reasonable yields of DNA from these clones. In addition, one cosmid clone (407-32.5G6) was unstable and had to be grown from the original frozen stock in order to obtain satisfactory DNA preparations.

10 The pWE15 vector allows the inserts to be excised with However, four Notl sites are present in the HVT genome, so that inserts spanning these sites cannot be excised with Notl. Two of the Notl sites are present in the BamHI #2 fragment of HVT, this fragment was The other two sites are 15 cloned directly in pSP64. present in the unique short region within the BamHI #1 fragment. This fragment was cloned directly in the The three sheared cosmids and the two pWE15 vector. BamHI fragments cover all but a small portion of the Because these regions are 20 ends of the HVT genome. repeated in the internal portions of the genome, all of the genetic information is available.

A StuI site within the HVT US2 gene was established as a useful site for foreign DNA insertion utilizing the HOMOLOGOUS RECOMBINATION PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUSES (see Example 6). The HVT US2 gene is located within the BamHI #1 fragment which contains five StuI sites. To facilitate the use of this site for insertion of foreign DNA by the StuI site within the US2 gene was converted to a unique HindIII site. This was accomplished by partially digesting the BamHI #1 subclone with StuI, and then inserting a 10 kb fragment conferring kanomycin resistance (Neo<sup>®</sup>) into the site using HindIII linkers. The kanomycin

resistance gene allowed positive selection of recombinant clones. The Neo<sup>2</sup> fragment was removed by digestion with *HindIII* followed by religation generating clone 430-84.215.

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DNA was prepared for reconstruction experiments by restriction digestion with enzymes which cut subclones outside or flanking the HVT insertions. some instances, one cosmid in a reconstruction was used undigested. Digested DNAs were extracted once with phenol precipitated with ethanol. DNA resuspended at a concentration of 0.5 to 1 ug/ml. Viral reconstruction experiments were performed Lipofectin (BRL) to mediate transfection. Two to three micrograms each of subclone were added to 0.5 ml of MEM (Earle's salts) supplemented with essential amino acids and 2% penicillin/Streptomysin (MEM+). Controls consisted of MEM+ with no DNA, with several ug of HVT DNA, or with 4 out of 5 of the Separately, 30  $\mu$ l of the Lipofectin were added to another 0.5 ml. of MEM+. These two mixtures were then combined and incubated at RT for 15 minutes.

Chick embryo fibroblast (CEF) cells were prepared for transfection in the following manner. CEFs (Spafas) were grown in 6 well dishes at 39°C in F10/M199 (1:1) media containing 1% non-essential amino acids, 2% penicillin/streptomycin, and 5% fetal calf serum (CEF+). Cells were transfected at a confluence of 90 - 95%. For transfection, wells were aspirated and rinsed 3 times with MEM+, and then incubated 4 hours at 39°C with the 1 ml lipofectin/DNA mixture above. One ml more of CEF+ was then added to the wells, and cells were incubated overnight and fed with CEF+. Plates were then examined daily for the appearance of plaques.

Lipofectin with control HVT DNA resulted in the

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appearance of plaques within 5 days. When only four of the five subclones were used, no plaques were obtained. When the five overlapping genomic fragments of HVT were to reconstruct the virus, plaques appeared anywhere from 5 to 19 days after the initial lipofection. In the case of plaques appearing late, plaques were not initially seen on the infected and it was only after passaging the monolayer, monolayer and replating on a larger surface that plaques appeared. After passaging, plaques generally appeared within 3 days. Recombinant viruses were plaque purified approximately three and then analyzed for insertion of foreign DNAs.

15 BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS. When the foreign gene encoded the enzyme  $\beta$ -galactosidase, the plaques that contained the gene were visualized more easily. The chemical Bluogal™ (Bethesda Research Labs) was incorporated at the level of 200-300  $\mu$ g/ml into the 20 agarose overlay during the plaque assay, and the plaques that expressed active  $\beta$ -galactosidase turned The blue plaques were then picked and purified by further blue plaque isolations. Other foreign genes were inserted by homologous recombination such that 25 they replaced the  $\beta$ -galactosidase gene; in instance non-blue plaques were picked for purification of the recombinant virus.

USING BLACK PLAQUE ASSAYS. To analyze expression of foreign antigens expressed by recombinant HVT viruses, monolayers of CEF cells are infected with recombinant HVT, overlaid with nutrient agarose media and incubated for 4-5 days at 39°C. Once plaques have developed, the agarose overlay is removed from the dish, the monolayer rinsed 1x with PBS, fixed with 100% methanol for 10 minutes at room temperature and the cells air dried.

After re-hydrating the plate with PBS, the primary antibody is diluted to the appropriate dilution with PBS and incubated with the cell monolayer for 2 hours to overnight at room temperature. Unbound antibody is then removed from the cells by washing three times with PBS at room temperature. An alkaline phosphatase conjugated secondary antibody is diluted with PBS and incubated with the cells for 2 hours temperature. Unbound secondary antibody is then removed by washing the cells three times with PBS at room temperature. Next, the monolayer is rinsed in color development buffer (100mM Tris pH 9.5/ 100mM NaCl/ 5mM MgCl2), and then incubated 10 minutes to overnight at room temperature with freshly prepared substrate solution (0.3 mg/ml Nitro Blue tetrazolium + 0.15 mg/ml 5-Bromo-4-Chloro-3-Indolyl Phosphatase in color development buffer.) Finally, the reaction is stopped by replacing the substrate solution with TE (10mM Tris, pH7.5/ 1 mM EDTA). Plaques expressing the correct antigen will stain black.

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PLAQUE HYBRIDIZATION PROCEDURE FOR ASSESSING THE PURITY OF RECOMBINANT HVT STOCKS. When no suitable immunological reagent exists to detect the presence of a particular antigen in a recombinant HVT virus, plaque hybridization can be used to assess the purity of a Initially, CEF cell monolayers are infected with various dilutions of the viral stocks to give ~50-100 plaques/10 cm.dish, overlaid with nutrient agarose media and incubated for 4-5 days at 39°C. Once plaque development occurs, the position of each plaque is marked on bottom of the dish. The agarose overlay is then removed, the plate washed with PBS, and the remaining CEF monolayer is transferred to a NC membrane or BioRad nylon membrane pre-wetted with PBS note of the membrane position relative to the dish). Cells contained on the NC membranes are then lysed by

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placing the membranes in 1.5 mls of 1.5M NaCl and 0.5M NaOH for five minutes. The membranes are neutralized by placing them in 1.5 mls of 3M Sodium acetate (pH 5.2) for five minutes. DNA from the lysed cells is then bound to the NC membranes by baking at 80°C for After this period the membranes are prehybridized in a solution containing 6X SSC, 3% skim milk, 0.5% SDS, ( $\pm$ ) salmon sperm DNA (50  $\mu$ g/ml) for one hour at 65°C. Radio-labeled probe DNA (alpha 32P-dCTP) is then added and the membranes incubated at 65°C overnight (~12 hours). After hybridization the NC membranes are washed two times (30 minutes each) with 2X SSC at 65°C, followed by two additional washes at 65°C with 0.5X SSC. The NC membranes are then dried and exposed to X-ray film (Kodak X-OMAT, AR) at -70°C for 12 hours. Positive signals are then aligned with the position of the plaques on the dish and purity of the stock is recorded as the percentage of positive plaques over the total.

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CONSTRUCTION OF HOMOLOGY VECTOR FOR INSERTION OF THE BETA-GALACTOSIDASE GENE INTO HVT US2 GENE. The betagalactosidase (lacZ) gene was inserted into the HVT EcoRI # 7 fragment at the unique StuI site. The marker gene is oriented in the same direction as the US2 gene. A detailed description of the marker gene is given in Figures 7A and 7B. It is constructed utilizing standard recombinant DNA techniques (Maniatis et al. 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources with the synthetic DNA sequences indicated in Figures 7A and 7B. Fragment 1 is an approximately 413 base pair SalI to BamHI restriction sub-fragment of the PRV BamHI restriction fragment 10 (Lomniczi et al., 1984). Fragment 2 is an approximately 3010 base pair BamHI to PvuII restriction fragment of plasmid pJF751 (Ferrari et al., 1985). Fragment 3 is an approximately 754 base pair NdeI to

SalI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi et al., 1984).

RNA ISOLATED FROM CONCANAVALIN A STIMULATED CHICKEN SPLEEN CELLS: Chicken spleens were dissected from 3 week old chicks from SPAFAS, Inc., washed, disrupted through a syringe/needle to release cells After allowing stroma and debri to settle out, cells were pelleted and washed twice with PBS. The cell pellet was treated with a hypotonic lysis buffer to lyse red blood cells, and splenocytes were recovered and washed twice with PBS. Splenocytes were resuspended at 5 x 106 cells/ml in RPMI containing 5% FBS and 5  $\mu$ g/ml Concanavalin A and incubated at 39° for 48 hours. Total RNA was isolated from the cells using guanidine isothionate lysis reagents and protocols from the Promega RNA isolation kit (Promega Corporation, Madison WI).  $4\mu g$  of total RNA was used in each 1st strand reaction containing the appropriate antisense primers and AMV reverse transcriptase (Promega Corporation, Madison WI). cDNA synthesis was performed in the same tube following the reverse transcriptase reaction, using the appropriate sense primers and Vent® DNA polymerase (Life Technologies, Inc. Bethesda, MD).

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SUBGENOMIC CLONE 172-07.BA2. Plasmid 172-07.BA2 was constructed for the purpose of generating recombinant HVT. It contains an approximately 25,000 base pair region of genomic HVT DNA. It may be used conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the following sources. The first fragment an

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approximately 2999 base pair BamHI to BamHI restriction fragment of pSP64 (Promega). The second fragment is the approximately 25,000 base pair BamHI #2 fragment of HVT (Buckmaster et al., 1988).

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HOMOLOGY VECTOR 172-29.31. The plasmid 172-29.31 was constructed for the purpose of inserting foreign DNA into HVT. It contains a unique XhoI restriction enzyme site into which foreign DNA may be inserted. plasmid containing a foreign DNA insert at the XhoI site is used according to the DNA COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES or the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS a virus containing the foreign DNA will result. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al. 1989), by joining restriction fragments from the following sources. The first fragment is an approximately 2999 base pair BamHI to BamHI restriction fragment of pSP64 (Promega). second fragment is the approximately 3300 base pair BamHI #16 fragment of HVT (Buckmaster et al., 1988). The complete sequence of the BamHI #16 fragment is given in SEQ ID NO:3. Note that the fragment was cloned the UL43 ORF is in the transcriptional orientation to the pSP64  $\beta$ -lacatamase gene.

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HOMOLOGY VECTOR 172-63.1. The plasmid 172-63.1 was constructed for the purpose of inserting foreign DNA into HVT. It contains a unique XhoI restriction enzyme site into which foreign DNA may be inserted. When a plasmid containing a foreign DNA insert at the XhoI site is used according to the DNA COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES or the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS a virus containing the foreign DNA

will result. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the following sources. The first fragment is an approximately 2999 base pair EcoRI to EcoRI restriction fragment of pSP64 (Promega). The second fragment is the approximately 5500 base pair EcoRI #9 fragment of HVT. Note that the EcoRI fragment was cloned such that the unique XhoI site is closest to the unique HindIII site in the pSP64 vector.

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HOMOLOGY VECTORS 255-18.B16. The plasmid 255-18.B16 was constructed for the purpose of inserting the NDV HN and F genes into HVT. The NDV HN and F genes were inserted as a SalI fragment into the homology vector 172-29.31 at the *XhoI* site. The NDV HN and F genes were inserted in the same transcriptional orientation the UL43 ORF in the parental homology vector. A detailed description of the SalI fragment is shown in Figures 12A-12C. The inserted SalI fragment may be constructed utilizing standard recombinant techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources with the synthetic DNA sequences indicated in Figures 12A, 12B and 12C. Fragment 1 is approximately 416 base pair SalI to restriction sub-fragment of the PRV BamHI restriction fragment 10 (Lomniczi et al., 1984). Fragment 2 is an approximately 3009 base pair BamHI to PvuII fragment of the plasmid pJF751 (Ferrari et al., 1985). Fragment 3 is an approximately 1200 base pair AvaII to EcoRI restriction fragment of full length NDV HN cDNA. Fragment 4 is an approximately 179 base pair EcoRI to restriction fragment of the plasmid pSP64 (Promega). Fragment 5 is an approximately 357 base pair Smal to BamHI restriction sub-fragment of the HSV-1 restriction fragment N. Fragment BamHI

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approximately 1812 base pair BamHI to PstI restriction fragment of the full length NDV F cDNA. Fragment 7 is an approximately 235 base pair PstI to ScaI restriction fragment of the plasmid pBR322.

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SUBGEMOMIC CLONE 378-50.BA1. Cosmid 378-50.BA1 was constructed for the purpose of generating recombinant HVT. It contains an approximately 29,500 base pair region of genomic HVT DNA. Ιt may be conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for construction of recombinant HVT. This cosmid may be constructed by joining two restriction fragments from the following sources. The first fragment is an approximately 8164 base pair BamHI to BamHI restriction fragment of pWE15 (Stratagene). The second fragment is the approximately 29,500 base pair BamHI #1 fragment of HVT (Buckmaster et al., 1988).

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SUBGEMOMIC CLONE 407-32.1C1. Cosmid 407-32.1C1 was constructed for the purpose of generating recombinant HVT. It contains an approximately 38,850 base pair region of genomic HVT DNA (see Figure 8). This region BamHI fragments 11, 7, 8, 21, 6, approximately 1250 base pairs of fragment 13, approximately 6,700 base pairs of fragment 1. It may be used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT. This cosmid maybe constructed as described above in the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. It was isolated from the sheared DNA library by screening with the probes P1 and P4 (described in Figure 8). A bacterial strain containing this cosmid has been deposited on March 3, 1993

pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. 75428.

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SUBGEMOMIC CLONE 407-32.2C3. Cosmid 407-32.2C3 was constructed for the purpose of generating recombinant HVT. It contains an approximately 40,170 base pair region of genomic HVT DNA (see Figure 8). This region includes BamHI fragments 10, 14, 19, 17, 5, approximately 2,100 base pairs of fragment 2. It may be used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT. This cosmid may be constructed as described above in the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. It was isolated from the sheared DNA library by screening with the probes P1 and P2 (described in Figure 8). A bacterial strain containing this cosmid has been deposited pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. 75430.

SUBGEMOMIC CLONE 407-32.5G6. Cosmid 407-32.5G6 was constructed for the purpose of generating recombinant HVT. It contains an approximately 40,000 base pair region of genomic HVT DNA (see Figure 8). This region includes BamHI fragments 9, 3, 20, 12, 16, 13, approximately 1,650 base pairs of fragment 2, and approximately 4,000 base pairs of fragment 11. It may be used in conjunction with other subgenomic clones

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according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT. This cosmid may be constructed as described above in the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. It was isolated from the sheared DNA library by screening with the probes P2 and P3 (described in Figure 8). A bacterial strain containing this cosmid has been deposited on March 3, 1993 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. 75427.

The plasmid 435-47.1 was HOMOLOGY VECTOR 435-47.1. constructed for the purpose of inserting foreign DNA It contains a unique HindIII restriction into HVT. enzyme site into which foreign DNA may be inserted. 20 When a plasmid containing a foreign DNA insert at the according to the HindIII site is used COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES or the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS virus SUBGENOMIC FRAGMENTS OVERLAPPING 25 containing the foreign DNA will result. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the The first fragment sources. 30 following approximately 2999 base pair EcoRI to EcoRI restriction The second fragment is fragment of pSP64 (Promega). the approximately 7300 base pair EcoRI #7 fragment of HVT. Note that the HindIII site of the pSP64 vector was removed by digesting the subclone with HindIII followed 35 by a Klenow fill in reaction and religation. A synthetic HindIII linker (CAAGCTTG) was then inserted

into the unique StuI site of the EcoRI #7 fragment.

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SUBGEMOMIC CLONE 437-26.24. Plasmid 437-26.24 constructed for the purpose of generating recombinant HVT. It contains an approximately 13,600 base pair region of genomic HVT DNA. It may be conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS construction of recombinant HVT. This plasmid may be constructed utilizing standard recombinant techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the following sources. The first fragment approximately 2970 base pair HindIII to BamHI restriction fragment of pSP64 (Promega). The second fragment is the approximately 13,600 base pair BamHI to StuI sub-fragment of the BamHI #2 fragment of HVT (Buckmaster et al., 1988). Note that the BamHI #2 fragment contains five StuI sites, the site utilized in this subcloning was converted to a HindIII site as described in the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS.

25 SUBGEMOMIC CLONE 437-26.26. Plasmid 437-26.26 was constructed for the purpose of generating recombinant HVT. It contains an approximately 15,300 base pair region of genomic HVT DNA. It may be used conjunction with other subgenomic clones according to 30 the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS construction of recombinant HVT. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the 35 following sources. The first fragment is an approximately 2970 base pair HindIII to BamHI

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restriction fragment of pSP64 (Promega). The second fragment is the approximately 15,300 base pair BamHI to StuI sub-fragment of the BamHI #2 fragment of HVT (Buckmaster et al., 1988). Note that the BamHI #2 fragment contains five StuI sites, the site utilized in this subcloning was converted to a HindIII site as described in the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS.

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10 HOMOLOGY VECTORS 456-18.18 and 456-17.22. The plasmids 456-18.18 and 456-17.22 were constructed for the purpose of inserting the MDV gA and gB genes into HVT. The MDV genes were inserted as a cassette into the homology vector 435-47.1 at the unique HindIII site. 15 The MDV genes were inserted at the blunt ended HindIII site as a blunt ended PstI to EcoRI fragment (see Figures 10A and 10B). The HindIII and EcoRI sites were blunted by the Klenow fill in reaction. The PstI site was blunted by the T4 DNA polymerase reaction. Note 20 that the MDV cassette was inserted in both orientations. Plasmid 456-18.18 contains the MDV genes inserted in the opposite transcriptional orientation to the US2 gene in the parental homology vector. Plasmid 456-17.22 contains the MDV genes inserted in the same 25 transcriptional orientation as the US2 gene in the parental homology vector. A detailed description of the MDV cassette is given in Figures 10A and 10B. may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 30 1989), by joining restriction fragments following sources with the synthetic DNA sequences indicated in Figures 10A and 10B. Fragment 1 is an approximately 2178 base pair PvuII to EcoRV restriction sub-fragment of the MDV **Eco**RI 6.9 KB genomic 35 restriction fragment (Ihara et al., 1989). Fragment 2 is an approximately 3898 base pair SalI to EcoRI genomic MDV fragment (Ross, et al., 1989).

HOMOLOGY VECTOR 528-03.37. The plasmid 528-03.37 was constructed for the purpose of inserting the infectious laryngotracheitis (ILT) virus gD gene into HVT. The gD gene followed by the PRV gX poly adenylation signal was inserted as a cassette into the homology vector 435-47.1 at the unique HindIII site. The cassette may be constructed utilizing standard recombinant techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments following sources. The fragment first is an approximately 2060 base pair EcoRI to BclI restriction sub-fragment of the ILT KpnI genomic restriction fragment #8 (10.6 KB). The second fragment is an approximately 754 base pair NdeI to SalI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi et al., 1984). Note that the fragments are oriented such that BclI and NdeI sites are contiguous.

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20 HOMOLOGY VECTOR 528-11.43. The plasmid 528-11.43 was constructed for the purpose of inserting the infectious laryngotracheitis (ILT) virus gB gene (A.M. Grifin, 1991) into HVT. The gB gene was inserted as an EcoRI fragment into the homology vector 435-47.1 at the 25 unique HindIII site. The gB gene was inserted at the blunt ended HindIII site as a blunt ended EcoRI fragment. The HindIII and EcoRI sites were blunted by the Klenow fill in reaction. The gB gene was inserted in the same transcriptional orientation as the US2 gene 30 in the parental homology vector. The EcoRI fragment may be obtained as a 3.0 KB ILT virus genomic fragment.

HOMOLOGY VECTOR 518-46.B3. The plasmid 518-46.B3 was constructed for the purpose of inserting foreign DNA into HVT. It contains a unique *HindIII* restriction enzyme site into which foreign DNA may be inserted. When a plasmid containing a foreign DNA insert at the

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HindIII site is used according to the DNA COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES or the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC **FRAGMENTS** virus containing the foreign DNA will result. This plasmid may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining three restriction fragments from the following sources. The first fragment approximately 1649 base pair PvuI to SalI restriction fragment of pSP64 (Promega). The second fragment is an approximately 1368 base pair PvuI to SalI restriction fragment of pSP65 (Promega). The third fragment is the approximately 3400 base pair XhoI to XhoI fragment of plasmid 437-47.1.

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The plasmid 535-70.3 was HOMOLOGY VECTOR 535-70.3. constructed for the purpose of inserting the MDV qB, and gA genes and the NDV F gene into HVT. The F gene inserted as a cassette into homology vector 456-17.22 at the HindIII site located between the MDV gA and gB genes (see Junction B, Figure 10A). The F gene under the control of the HCMV immediate early followed by the promoter and HSV-1 TK adenylation signal. The F gene was inserted in the same transcriptional orientation as the US2 gene in the parental homology vector. The cassette may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from following sources. The first fragment approximately 1191 base pair PstI to AvaII restriction sub-fragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The second fragment is an approximately 1812 base pair BamHI to PstI restriction fragment of the full length NDV F cDNA clone strain). The last fragment is an approximately 784 base

pair SmaI to SmaI restriction sub-fragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

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HOMOLOGY VECTOR 549-24.15. The plasmid 549-24.15 was constructed for the purpose of inserting the MDV gB, and gA genes and the NDV HN and F genes into HVT. The HN and F genes were inserted as a cassette into homolgy vector 456-17.22 at the HindIII site located between the MDV gA and gB genes (see Junction B, Figure 10A). The HN and F genes are under the control of the PRV gpX and HCMV immediate early promoters respectively. The HN and F genes are followed by the poly and HSV-1 TK adenylation respectively. The cassette may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources. The first fragment is an approximately 413 base pair SalI to BamHI restriction sub-fragment of the fragment #10 (Lomniczi, et al., 1984) The second fragment is an approximately 1811 base pair AvaII to Nael restriction fragment of the full length NDV HN cDNA clone (B1 strain). The third fragment is approximately 754 base pair NdeI to SalI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi, et al., 1984). The fourth fragment is an approximately 1191 base pair PstI to AvaII restriction sub-fragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The fifth fragment is an approximately 1812 base pair BamHI to PstI restriction fragment of the full length NDV F cDNA clone (B1 strain). The last fragment is an approximately 784 base pair SmaI to SmaI restriction sub-fragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

HOMOLOGY VECTOR 549-62.10. The plasmid 549-62.10 was constructed for the purpose of inserting the MDV gB,

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and gA genes and the NDV HN gene into HVT. The HN gene inserted as a cassette into homolgy vector 456-17.22 at the HindIII site located between the MDV qA and gB genes (see Junction B, Figure 10A). The HN gene under the control of the PRV gpX promoter and followed by the PRV gX poly adenylation signal. The HN gene was inserted in the same transcriptional orientation as the US2 gene in the parental homology The cassette may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources. The fragment is an approximately 413 base pair SalI to restriction sub-fragment of the PRV fragment #10 (Lomniczi, et al., 1984) The second fragment is an approximately 1811 base pair AvaII. to Nael restriction fragment of the full length NDV HN cDNA clone (B1 strain). The last fragment approximately 754 base pair NdeI to SalI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi, et al., 1984).

550-60.6. SUBGENOMIC CLONE Plasmid 550-60.6 constructed for the purpose of generating recombinant HVT. It contains an approximately 12,300 base pair region of genomic HVT DNA. It may be used conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS OVERLAPPING SUBGENOMIC FRAGMENTS FROM for construction of recombinant HVT. This plasmid may be standard constructed utilizing recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining two restriction fragments from the following The first sources. fragment approximately 4176 base pair EcoRV to BamHI restriction fragment of pBR322. The second fragment approximately 12,300 base pair sub-fragment of the 5

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BamHI #2 fragment of HVT (Buckmaster et al., 1988). This fragment was generated in the following manner. Plasmid 437-26.26 was linearized with HindIII and then resected with the ExoIII Mung Bean Deletion Kit (Stratagene). Samples from the 3 and 4 minute reactions were combined and digested with BamHI resulting in a population of fragments containing the desired 12,300 base pair sub-fragment. This population was cloned into the pBR322 fragment and the resulting clones were screened for the appropriate size and restriction map. Fortuitously the resected sub-fragment that generated clone 550-60.6 ended in the nucleotides GG which generated a second BamHI site when ligated to the EcoRV site (ATCC) of pBR322. A bacterial strain containing this plasmid has been deposited on March 3, pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. 75429.

HOMOLOGY VECTORS 566-41.5. The plasmid 566-41.5 was constructed for the purpose of inserting the MDV qA, qB and gD genes into HVT. The MDV gD gene was inserted as a HindIII fragment into the homology vector 456-17.22 at the HindIII site located between MDV gA and gB (see Figures 10A and 10B). The MDV gene was inserted in the same transcriptional orientation as gA and gB in the parental homology vector. A detailed description of the HindIII fragment containing the MDV qD gene is shown in Figures 11A and 11B. Note that herpesvirus а polyadenation signal was added to the qD gene cassette. The inserted HindIII fragment may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources with

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the synthetic DNA sequences indicated in Figures 11A and 11B. Fragment 1 is an approximately 784 base pair SmaI to SmaI restriction sub-fragment of the HSV-1 BamHI restriction fragment Q (McGeoch et al., 1988). Note that this fragment is oriented such that the polyadenylation sequence (AATAAA) is located closest to junction B. Fragment 2 is an approximately 2177 base pair SalI to NcoI sub-fragment of the MDV BglII 4.2 KB genomic restriction fragment (Ross, et al., 1991).

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HOMOLOGY VECTOR 567-72.1D. The plasmid 567-72.1D was constructed for the purpose of inserting the MDV gB, qA, and qD genes and the infectious bronchitis virus (IBV) matrix and spike genes into HVT. The IBV genes inserted as a cassette into homolgy vector 566-41.5 at the unique NotI site located upstream of the MDV gD gene (see Junction C, Figure 11B). spike and matrix genes are under the control of the **HCMV** immediate early and PRV Xqp promoters respectively. The IBV spike and matrix genes are followed by the HSV-1 TK and PRV gX poly adenylation signals respectively. The IBV genes were inserted in the same transcriptional orientation as the US2 gene in the parental homology vector. The cassette may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources. The first fragment is an approximately 413 base pair SalI to BamHI restriction sub-fragment of the PRV BamHI fragment #10 (Lomniczi, et al., 1984) The second fragment contains amino acids 1 to 223 of the IBV matrix gene. The coding region was obtained from a cDNA clone of the Arkansas strain of IBV. The third fragment is an approximately 754 base pair NdeI to SalI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi, et al., 1984). The fourth fragment is an approximately 1191 base pair

PstI to AvaII restriction sub-fragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The fifth fragment contains amino acids 4 to 1162 of the IBV spike gene. The coding region was obtained from a cDNA clone of the Arkansas strain of IBV. The last fragment is an approximately 784 base pair SmaI to SmaI restriction sub-fragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

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10 HOMOLOGY VECTOR 603-57.F1. The plasmid 603-57.F1 was constructed for the purpose of inserting the IBDV VP2 gene into HVT. The IBDV VP2 gene was inserted as a cassette into homolgy vector 435-47.1 at the unique HindIII site. The VP2 gene is under the control of the 15 HCMV immediate early promoter and is followed by the HSV-1 TK poly adenylation signal. The VP2 gene was inserted in the same transcriptional orientation as the US2 in the parental homology vector. The cassette may be constructed utilizing standard recombinant 20 techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources. The first fragment approximately 1191 base pair PstI to AvaII restriction sub-fragment of the HCMV genomic XbaI E fragment (D.R. 25 Thomsen, et al., 1981). The second fragment is an approximately 1081 base pair BclI to BamHI restriction sub-fragment of the full length IBDV cDNA clone (see SEQ ID NO:1). Note that the BclI site was introduced into the cDNA clone directly upstream of the VP2 30 initiator methionine by converting the sequence CGCAGC to TGATCA. The first and second fragments are oriented such that AvaII and BclI sites are contiguous. The third fragment is an approximately 784 base pair SmaI to Smal restriction sub-fragment of the HSV-1 BamHI 35 restriction fragment Q (McGeoch, et al., 1985).

HOMOLOGY VECTOR 633-13.27. The plasmid 633-13.27 was

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constructed for the purpose of inserting the MDV gB, gA and gD genes and the NDV HN and F genes into HVT. The HN and F genes are under the control of the PRV gpX and HCMV immediate early promoters respectively. The HN and F genes are followed by the PRV gX poly and HSV-1 TK adenylation signals respectively. All five genes were inserted in the same transcriptional orientation as the US2 gene in the parental homology vector. The genes were inserted in the following order MDV gA, NDV HN, NDV F, MDV gD, and MDV qB.

HOMOLOGY VECTOR 634-29.16. The plasmid 634-29.16 was constructed for the purpose of inserting the ILT virus gB and gD genes into HVT. The lacZ marker gene followed by the ILT gB and gD genes inserted as a cassette into the homology vector 172-29.31 at the unique XhoI site. The cassette may be constructed utilizing standard recombinant DNA techniques (Maniatis et al, 1982 and Sambrook et al, 1989), by joining restriction fragments from the following sources. The first fragment is an approximately 4229 base pair Sall to Sall restriction fragment derived from the lacZ marker gene described above and shown in Figures 7A and 7B. The second fragment is an approximately 2060 base pair EcoRI to BclI restriction sub-fragment of the ILT KpnI restriction fragment #8 (10.6 KB). The third fragment an approximately 754 base pair NdeI restriction sub-fragment of the PRV BamHI restriction fragment #7 (Lomniczi et al., 1984). Note that the second and third fragments are oriented such that BclI and NdeI sites are contiguous. The fourth fragment is the 3.0 KB ILT virus genomic EcoRI fragment containing gene. All three genes are in the transcriptional orientation as the UL43 gene.

SUBGENOMIC CLONE 415-09.BA1. Cosmid 415-09.BA1 was constructed for the purpose of generating recombinant

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It contains an approximately 29,500 base pair HVT. BamHI #1 fragment of genomic HVT DNA. It was used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT. This cosmid was constructed by joining two restriction (Sambrook, et al., 1989) from the following sources. The vector is an approximately 4430 base pair BamHI to BamHI restriction fragment of pSY1005 derived from pHC79 (Bethesda Research Labs, Inc.) and pWE15 (Stratagene, Inc.). The first fragment is approximately 29,500 base pair BamHI #1 fragment of the HVT genome (Buckmaster et al., 1988).

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SUBGENOMIC CLONE 672-01.A40. Cosmid 672-01.A40 was constructed for the purpose of generating recombinant HVT. It was isolated as a subclone of cosmid 407-32.1C1 (see Figures 8 and 15). Cosmid 672-01.A40 contains an approximately 14,000 base pair NotI to AscI subfragment and an approximately 1300 base pair AscI to BamHI subfragment of cosmid 407-32.1C1. The cosmid was constructed by joining restriction fragments (Sambrook, et al., 1989) from the following sources. The vector is an approximately 2700 base pair NotI to BamHI fragment constructed from pNEB193 (New England Biolabs, Inc.) which contains a NotI linker inserted into the SmaI site. Fragment 1 is an approximately 15,300 base pair region of genomic HVT DNA. This region includes BamHI fragments 11 and 7, and approximately 1250 base paris of fragment 13. It was used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction ofrecombinant HVT.

SUBGENOMIC CLONE 654-45.1. Plasmid 654-45.1 was

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constructed for the purpose of generating recombinant HVT. It was isolated as an AscI subclone of cosmid 407-(see Figures 8 and 15). The cosmid was constructed by joining restriction fragments (Sambrook, et al., 1989) from the following sources. is an approximately 2000 base pair AscI fragment constructed from a 2000 base pair AatII to PvuII fragment of pNEB 193 (New England Bilabs, Inc.) blunt ended with Klenow DNA polymerase and AscI linkers Fragment 1 is an approximately 8600 base pair AscI to AscI fragment of genomic HVT DNA. This region includes BamHI fragments 10 and 21. approximately 1100 base pairs of fragment 6 approximately 1300 base pairs of fragment 7. The XhoI site (Nucleotide #1339-1344; SEO ID NO. 48) has been converted to a unique PacI site using synthetic DNA The PacI site was used in insertion and expression of foreign genes in HVT. (See Figure 13A). It was used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT.

SUBGENOMIC CLONE 686-63.A1. Plasmid 686-63.A1 was constructed for the purpose of generating recombinant It was isolated as an AscI subclone of cosmid (see Figure 8, 15). The cosmid was 407-32.1C1 constructed by joining restriction fragments (Sambrooks, et al., 1989) from the following sources. The vector is an approximately 2000 base pair AscI fragment constructed from a 2000 base pair AatII to PvuII fragment of pNEB193 (New England Biolabs, Inc.) blunt ended with Klenow DNA polymerase and AscI linkers inserted. Fragment 1 is an approximately 8600 base pair AscI to AscI fragment of genomic HVT DNa. This region includes BamHI fragments 10 and 21, and approximately 1100 base pairs of fragment 6

approximately 1300 base pairs of fragment 7. The XhoI site (Nucleotide #1339-1344; SEQ ID NO. 48) has beenconverted to a unique NotI site using synthetic DNA linkers. The NotI site was used for the insertion and expression of foreign genes in HVT. (See Figure 13B). It was used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT.

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SUBGENOMIC CLONE 672-07.C40. Cosmid 672-07.C40 was constructed for the purpose of generating recombinant HVT. It was isolated as a subclone of cosmid 407-32.1C1 (see Figures 8 and 15). Cosmid 672-07.C40 contains an approximately 1100 base pair BamHI to AscI subfragment and an approximately 13,000 base pair AscI to NotI subfragment of cosmid 407-32.1C1. The cosmid was constructed by joining restriction fragments (Sambrook, et al., 1989) from the following sources. The vector is an approximately 2700 base pair NotI to BamHI fragment constructed from pNEB193 ( New England Biolabs, Inc.) which contains a NotI linker inserted into the Smal site. Fragment 1 is an approximately 14,100 base pair region of genomic HVT DNA. This region includes BamHI fragments 6 and 18, and an approximately 2600 base pair BamHI to NotI fragment within BamHI fragment #1. It was used in conjunction with other subgenomic clones according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS for the construction of recombinant HVT.

SUBGENOMIC CLONE 706-57.A3. Plasmid 706-57.A3 was constructed for the purpose of generating recombinant HVT. Plasmid 706-57.A3 contains the IBDV VP2 gene inserted into the *PacI* site of plasmid 654-45.1. The IBDV VP2 gene uses the IBRV VP8 promoter and ILTV US3 polyadenylation signal. The cosmid was constructed

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utilizing standard recombinant DNA techniques (Sambrook, et al., 1989). The first fragment is a 208 base pair HindIII to BamHI fragment coding for the IBRV VP8 promoter (Carpenter, et al., 1991)). The second fragment is an approximately 1626 base pair fragment coding for the IBDV VP2 gene derived by reverse transcription and polymerase chain reaction (Sambrook, et al., 1989) of IBDV standard challenge strain (USDA) genomic RNA (Kibenge, et al., 1990). The antisense primer used for reverse transcription and PCR was 5'-CTGGTTCGGCCCATGATCAGATGACAAACCTGCAAGATC-3' (SEQ ID NO. 53). The sense primer used for PCR CTGGTTCGGCCCATGATCAGATGACAAACCTGCAAGATC-3' (SEQ ID NO. The DNA fragment generated by PCR was cloned into the PCR-Direct $^{\text{TM}}$  vector (Clontech Laboratories, Inc., Pali Alto, CA). The IBDV VP2 fragment was subcloned next tot he VP8 promoter using BclI sites generated by the PCR primers. The DNA sequence at this junction adds amino acids methionine, aspartate and glutamine before the antive initiator methionine of VP2. The DNA fragment contains the coding sequence from amino acid 1 to amino acid 536 of the IBDV polyprotein (SEQ ID NO: 2) which includes the entire coding sequence of the VP2 The third fragment is an approximately 494 protein. base pair fragment coding for the ILTV US3 polyadenylation signal.

SUBGENOMIC CLONE 711-92.1A. Plasmid 711-92.1A was constructed for the purpose of generating recombinant HVT. Plasmid 711-92.1A contains the ILTV gD and gI genes inserted into the PacI site of plasmid 654-45.1. qD and gI genes use their respective endogenous ILTV promoters and single shared endogenous polyadenylation signal. The plasmid was constructed utilizing standard' recombinant DNA techniques (Sambrook, et al., 1989). The first fragment is an approximately 3556 base pair SalI to HindIII

restriction subfragment of the ILTV Asp718I genomic fragment #8 (10.6 kb).

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**SUBGENOMIC CLONE 717-38.12**. Plasmid 717-38.12 constructed for the purpose of generating recombinant HVT. Plasmid 717-38.12 contains the NDV HN and F genes inserted into the PacI site of plasmid 654-45.1. The NDV HN gene uses the PRV gX promoter and the PRV gX polyadenylation signal. The NDV F gene uses the HCMV immediate early promoter and the HSV TK polyadenylation signal. The plamid was constructed utilizing standard recombinant DNA techniques (Sambrook, et al., 1989). The first fragment is an approximately 413 base pair SalI to BamHI restriction subfragment of the PRV BamHI fragment #10 (Lomniczi, et al., 1984). fragment is an approximately 1811 base pair AvaII to Nael restriction fragment of the full length NDV HN cDNA clone (B1 strain). The third fragment is an approximately 754 base pair NdeI to SalI restriction subfragment of the PRV BamHI restriction fragment #7 (Lomniczi, et al., 1984). The fourth fragment is an approximately 1191 base pair PstI to AvaII restriction subfragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The fifth fragment is an approximately 1812 base pair BamHI to PstI restriction fragment of the full length NDV F cDNA clone (B1 strain; SEQ ID NO: 12). The sixth fragment is an approximately 784 base pair SmaI to SmaI restriction subfragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

SUBGENOMIC CLONE 721-38.1J. Cosmid 721-38.1J was constructed for the purpose of inserting the MDV gA, gD, and gB genes into the unique short of HVT and for the purpose of generating recombinant HVT. Cosmid 721-38.1J contains the MDV gA, gD and gB genes inserted into a StuI site in the HVT US2 gene converted to a

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unique HindIII site within the BamHI #1 fragment of the unique short region of HVT. This region of the HVT BamHI #1 fragment containing the MDV genes was derived from S-HVT-062. Cosmid 721-38.1J was constructed by a partial restriction digest with BamHI of S-HVT-062 DNA and isolation of an approximately 39,300 base pair fragment. The cosmid was constructed utilizing standard recombinant DNA techniques (Sambrook, et al., joining restriction fragments from following sources. The vector is an approximately 8200 base pair BamHI fragment from cosmid vector pWE15. The first fragment is an approximately 900 base pair BamHI fragment from the repeat region of the HVT genome. The second fragment is an approximately 15,500 base pair BamHI to StuI subfragment of BamHI #1 of HVT. The third fragment is an approximately 8400 base pair cassette containing the MDV gA, gD, and gB genes (see figures 10 and 11). The fourth fragment is an approximately 14,500 base pair HindIII to BamHI subfragment of the BamHI #1 of HVT.

SUBGENOMIC CLONE 722-60.E2. Cosmid 722-60.E2 was constructed for the purpose of inserting the MDV gA, gD, and gB genes and the NDV HN and F genes into the unique short of HVT and for the purpose of generating recombinant HVT. Cosmid 722-60.E2 contains the MDV qA, gD and gB genes and the NDV HN and F genes inserted into a StuI site in the HVT US2 gene converted to a unique HindIII site within the BamHI #1 fragment of the unique short region of HVT. All five genes were inserted in the same transcriptional orientation as the HVT US2 gene. This region of the HVT BamHI #1 fragment containing the MDV and NDV genes was derived from S-HVT-106. Cosmid 722-60.E2 was constructed by a partial S-HVT-106 restriction digest with BamHI of and isolation of an approximately 46,300 base pari fragment. The cosmid was constructed utilizing

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standard recombinant DNA techniques (Sambrook, et al., 1989) by joining restriction fragments from following sources. The vector is an approximately 6100 base pair BamHI fragment from cosmid vector pSY1626 derived from pHC79 (Bethesda Research Labs, Inc.) and pWE15 (Strategene, Inc.). The first fragment is an approximately 900 base pair BamHI fragment from the repeat region of the HVT genome. The second fragment is approximately 15,500 base pair BamHI subfragment of BamHI #1 of HVT. The third fragment is an approximately 15,400 base pair cassette containing the MDV qA gene, (Figures 10A and 10B, SEQ ID NO: 8), the PRV gX promoter (Lomniczi et al., 1984), the NDV HN gene (SEQ ID NO: 10), the PRV gX polyadenylation site (Lomniczi et al., 1984), the HCMV immediate early promoter (D.R. Thomsen, et al., 1981), the NDV F gene (SEQ ID NO: 12), the HSV TK polyadenylation site (McGeoch, et al., 1985), the MDV gD gene (Figures 11A and 11B), the approximately 450 base pair ILTV US3 polyadenylation site, and the MDV gB gene (Figures 10A and 10B). The fourth fragment is an approximately 14,500 base pair StuI to BamHI subfragment of the BamHI #1 of HVT.

25 SUBGENOMIC CLONE 729-37.1. Plasmid 729-37.1 constructed for the purpose of generating recombinant HVT. Plasmid 729-37.1 contains the ILTV gD and gB genes inserted into the NotI site of plasmid 686-63.Al. The ILTV gD and gB genes use their respective endogenous 30 ILTV promoters, and the ILTV gD and gB gene are each followed by a PRV gX polyadenylation signals. The ILTV gD and gB gene cassette was constructed utilizing standard recombinant DNA techniques (Sambrook, et al., 1989). The first fragment is an approximately 2052 base 35 pair SalI to XbaI restriction subfragment of the ILTV Asp718I genomic fragment #8 (10.6 kb). The second fragment is an approximately 572 base pair XbaI to

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Asp718I restriction subfragment of the PRV BamHI restriction fragment #7 (Lomniczi et al., 1984). The third fragment is an approximately 3059 base pair EcoRI to EcoRI restriction fragment of ILTV genomic DNA. The fourth fragment is an approximately 222 base pair EcoRI to SalI restriction subfragment of the PRV BamHI restriction fragment #7 (Lomniczi et al., 1984).

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SUBGENOMIC CLONE 739-27.16. Cosmid 739-27.16 constructed for the purpose of constructing achimeric HVT/MDV virus containing the HVT genes of the unique long region and the MDV type 1 genes of the unique short region. Cosmid 739-27.16 contains the complete unique short region of MDV type 1. This region contians the entire Smal B fragment and two Smal K fragments. Cosmid 739-27.16 was constructed by a partial restriction digest with SmaI of MDV DNA and isolation of an approximately 29,000 to 33,000 base pair fragment. The cosmid was constructed utilizing standard recombinant DNA techniques (Sambrook, et al., by joining restriction fragments from following sources. The vector is an approximately 8200 base pair BamHI fragment (made blunt-ended with Lenow DNa polymerase) from cosmid vector pWE15. The first fragment is an approximately 4050 base pair SmaI K fragment from the short internal repeat region of the MDV genome. The second fragment is an approximately 21,000 base pair fragment Smal B of MDV. The third fragment is an approximately 3,650 base pair Smal K fragment from the short terminal repeat region of the MDV genome (Fukuchi, et al., 1984, 1985).

SUBGENOMIC CLONE 751-87.A8. Plasmid 751-87.A8 was constructed for the purpose of generating recombinant HVT. Plasmid 751-87.A8 contains the chicken myelomonocytic growth factor (cGMF) gene inserted into the PacI site of plasmid 654-45.1. The cMGF gene uses

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the HCMV immediate early promoter and HSV-1 polyadenylation signal. The cosmid was constructed standard recombinant DNA techniques utilizing (Sambrook, et al., 1989). The following fragments were inserted into the PacI site of HVT subgenomic clone 654-45.1. The first fragment is an approximately 1191 base pair PstI to AvaII restriction subfragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The second fragment is an approximately 640 base pair fragment coding for the cMGF gene (58) derived by reverse transcription and polymerase chain reaction (PCR) (Sambrook, et al., 1989) of RNA ISOLATED FROM CONCANAVALIN A STIMULATED CHICKEN SPLEEN CELLS. The antisense primer used for reverse transcription and PCR was 5'-CGCAGGATCCGGGGCGTCAGAGGCGGGGGGGGGTG-3' (SEQ ID 57). The sense primer used for PCR was NO: GAGCGGATCCTGCAGGAGAGACACAGAGCTG-3' (SEQ ID NO: 58). The cMGF fragment was subcloned next to the HCMV IE promoter using BamHI sites generated by the PCR primers. The DNA fragment contains the coding sequence from amino acid 1 to amino acid 201 of the cMGF protein (58) which includes a 23 amino acid leader sequence at the amino terminus and 178 amino acids of the mature cMGF protein. The third fragment is an approximately 784 base pair SmaI to SmaI restriction subfragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

SUBGENOMIC CLONE 761-07.A1. Plasmid 761-07.A1 was constructed for the purpose of generating recombinant HVT. Plasmid 761-07.A1 contains the chicken interferon gene inserted into the PacI site of plasmid 654-45.1. The chicken interferon gene uses the HCMV immediate early promoter and HSV-1 TK polyadenylation signal. The cosmid was constructed utilizing standard recombinant DNA techniques (Sambrook, et al., 1989). The following fragments were inserted into the PacI site of HVT

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subgenomic clone 654-45.1. The first fragment is an approximately 1191 base pair PstI to AvaII restriction subfragment of the HCMV genomic XbaI E fragment (D.R. Thomsen, et al., 1981). The second fragment is an approximately 577 base pair fragment coding for the chicken interferon gene (59) derived by reverse transcription and polymerase chain reaction (PCR) et al., (Sambrook, 1989) of RNA ISOLATED FROM CONCANAVALIN A STIMULATED CHICKEN SPLEEN CELLS. The antisense primer used for reverse transcription and PCR was 5'-TGTAGAGATCTGGCTAAGTGCGCGTGTTGCCTG-3' (SEQ ID NO: 59). The sense primer used for PCR was TGTACAGATCTCACCATGGCTGTGCCTGCAAGC-3' (SEQ ID NO: 60). The chicken interferon gene fragment was subcloned next to the HCMV IE promoter using BqlII sites generated by the PCR primers. The DNA fragment contains the coding sequence from amino acid 1 to amino acid 193 of the chicken interferon protein (59) which includes a 31 amino acid signal sequence at the amino terminus and 162 amino acids of the mature protein encoding chicken interferon. The third fragment is an approximately 784 base pair Smal to Smal restriction subfragment of the HSV-1 BamHI restriction fragment Q (McGeoch, et al., 1985).

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#### EXAMPLE 1

# S-HVT-001

5 S-HVT-001 is a herpesvirus of turkeys (HVT) contains the E. coli  $\beta$ -galactosidase gene inserted into the unique long region of the HVT genome. restriction enzyme map of HVT has been published (T. Igarashi, et al., 1985). This information was used as 10 a starting point to engineer the insertion of foreign genes into HVT. The BamHI restriction map of HVT is shown in Figure 1A. From this data, several different regions of HVT DNA into which insertions of foreign genes could be made were targeted. The foreign gene 15 chosen for insertion was the  $E.~coli~\beta$ -galactosidase (lacZ) gene , which was used in PRV. The promoter was the PRV gpX promoter. The lacZ gene was inserted into the unique long region of HVT, specifically into the XhoI site in the BamHI #16 (3329bp) fragment, and was 20 shown to be expressed in an HVT recombinant by the formation of blue plaques using the substrate Bluogal™ (Bethesda Research Labs). Similarly, the lacZ gene has been inserted into the Sall site in the repeat region contained within the BamHI #19 (900 bp) fragment.

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These experiments show that HVT is amenable to the procedures described within this application for the insertion and expression of foreign genes in herpesviruses. In particular, two sites for insertion of foreign DNA have been identified (Figs. 1B and 1C).

# EXAMPLE 2

#### S-HVT-003

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S-HVT-003 is a herpesvirus of turkeys (HVT) that contains the  $E.~coli~\beta$ -galactosidase (lacZ) gene and

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the infectious bursal disease virus (IBDV) strain S40747 large segment of RNA (as a cDNA copy) (SEQ ID NO: 1) inserted into the unique long region of the HVT This IBDV DNA contains one open reading frame that encodes three proteins (5'VP2-VP4-VP3 3') (SEQ ID NO: 2), two of which are antigens to provide protection against IBDV infections of chickens. Expression of the genes for both  $\beta$ -galactosidase and the IBDV polyprotein are under the control of the pseudorabies virus (PRV) qpX gene promoter. S-HVT-003 was made by homologous S-HVT-003 was deposited on July 21, recombination. the Budapest Treaty 1987 pursuant to International Deposit of Microorganism for Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR 2178.

IBDV genes were cloned by the cDNA CLONING The PROCEDURE. Clones representing the genome of IBDV were screened by SOUTHERN BLOTTING OF DNA procedure against blots containing authentic IBDV RNA. Positive clones were then characterized by restriction mapping to Two such clones were identify groups of clones. identified, that together were found to represent the entire coding region of the IBDV large segment of RNA One cDNA clone (2-84) contained an (3.3 kb dsRNA). approximately 2500 base pair fragment representing the The second clone (2-40) first half of the IBDV gene. contained an approximately 2000 base pair fragment representing the distal half of the IBDV gene. Plasmid 2-84/2-40, representing the entire IBDV gene, constructed by joining clone 2-84 and 2-40 at a unique PvuII site present in the overlapping sequences. IBDV genome can be obtained from plasmid 2-84/2-40 as an approximately 3400 base pair Smal to Hpal fragment. Confirmation of the nature of the proteins encoded by

the IBDV gene was obtained by expressing the clone (2-84/2-40) in E. coli and detecting VP3 antigen using antiserum made against purified IBDV capsid proteins on Western blots. The cDNA of the IBDV large segment of RNA encoding the IBDV antigens show one open reading frame that will henceforth be referred to as the IBDV gene. The sequence of an Australian IBDV strain has published which bears homology close applicants' sequence (Hudson et al, 1986). Comparison of the amino acid differences between the two viruses revealed 29 amino acid changes within the 1012 amino There were only 3 amino acid acid coding region. differences deduced for VP4 and only 8 in VP3. contrast, VP2 contained 18 amino acid changes, 14 of which were clustered between amino acids 139 to 332.

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For insertion into the genome of HVT, the coding region for the IBDV gene was cloned between the PRV gpX promoter and the HSV TK poly-A signal sequence, creating plasmid 191-23. To aid in the identification of HVT recombinants made by homologous recombination containing the IBDV gene, the gpX promoted IBDV fragment from plasmid 191-23 was inserted behind (in tandem to) a lacZ gene controlled by a gpX promoter. The resultant plasmid, 191-47, contains the E.coli lacZ gene and the IBDV gene under the control of individual PRV gpX promoters. In constructing plasmid 191-47, various DNA fragments were joined by recombinant DNA techniques using either naturally occurring restriction sites or synthetic linker DNA. Details concerning the construction of these genes contained in plasmid 191-47 can be seen in Figures 2A, 2B, 2C and 2D.

The first segment of DNA (Segment 1, Figure 2A) contains the gpX promoter region including the residues encoding the first seven amino acids of the gpX gene, and was derived from a subclone of the PRV BamHI #10

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fragment as an approximately 800 base pair SalI to BamHI fragment. The second segment of DNA (Segment 2, Figure 2A) contains the E. coli  $\beta$ -galactosidase coding region from amino acid 10 to amino acid 1024 and was 5 derived from the plasmid pJF751 (obtained from Jim Hoch, Scripps Clinic and Research Foundation) as an approximately 3300 base pair BamHI to BalI fragment followed by an approximately 40 base pair Ava I to Sma The third segment of DNA (Segment 3, 10 Figure 2A) contains the gpX poly A signal sequence and was derived from a subclone of the PRV BamHI fragment as an approximately 700 base pair NdeI to StuI fragment. Segment three was joined to segment two by ligating the NdeI end which had been filled 15 according to the POLYMERASE FILL-IN REACTION, Smal site. The fourth segment of DNA (Segment 4, Figure 2A) contains the qpX promoter (TATA box and cap site) and was derived from a subclone of the PRV BamHI #10 fragment as an approximately 330 base pair NaeI to 20 AluI fragment. Additionally, segment four contains approximately 36 base pairs of HSV TK 5'untranslated leader sequence as a PstI to BglII fragment in which the PstI site has been joined to the AluI site through the use of a synthetic DNA linker (McKnight and 25 DNA segments four through six were Kingbury, 1982). inserted as a unit into the unique Kpn I site of segment three which is located 3' of the gpX poly A signal sequence. The fifth segment of DNA (Segment 5, Figure 2A) contains the entire coding region of the 30 large segment of RNA (cDNA clone) approximately 3400 base pair SmaI to HpaI fragment. The Smal site of segment five was fused to the BglII site of segment four which had been filled in according to the POLYMERASE FILL IN REACTION. Expression of the 35 IBDV gene (5'VP2-VP4-VP3 3') is under the control of the gpX promoter (segment 4), but utilizes its own natural start and stop codons. The sixth segment of DNA

(Segment 6, Figure 2A) contains the HSV TK poly-A signal sequence as an approximately 800 base pair SmaI fragment (obtained from Bernard Roizman, Univ. of Chicago). The HpaI site of segment five was fused to the SmaI site of segment six through the use of a synthetic DNA linker.

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In summary, the construct used to create S-HVT-003 (plasmid 191-47) contains (5' to 3') the PRV promoter, the gpX TATA box, the gpX cap site, the first seven amino acids of qpX, the E. coli  $\beta$ -galactosidase (lacZ) gene, the PRV poly-A signal sequence, the PRV gpX promoter, the gpX TATA box, the gpX cap site, a fusion within the qpX untranslated 5' leader to the IBDV gene. IBDV start codon, a fusion within the IBDV untranslated 3' end to HSV TK untranslated 3' end, and the TK poly-A signal sequence. The cassette containing these genes was engineered such that it was flanked by two EcoRI restriction endonuclease sites. As a result, approximately 9100 base pair fragment containing both lacZ gene and the IBDV gene can be obtained by digestion with EcoRI. Henceforth, the 9161 base pair EcoRI fragment will be referred to as the IBDV/lacZ cassette. The following procedures were used to construct S-HVT-003 by homologous recombination. IBDV/lacZ cassette was inserted into the unique XhoI site present within a subclone of the HVT BamHI #16 To achieve this, the XhoI site was first changed to an EcoRI site through the use of an EcoRI This site had previously been shown to be nonessential in HVT by the insertion of lacZ (S-HVT-It was also shown that the flanking homology regions in BamHI #16 were efficient in homologous recombination. Shown in Figures 3A and 3B, the genomic location of the BamHI #16 fragment maps within the unique long region of HVT. The BamHI #16 fragment is approximately 3329 base pairs in length (SEQ ID NOs:

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3, 4, 5, 6, and 7). HVT DNA was prepared by the PREPARATION **HERPESVIRUS** OF DNA procedure. Cotransfections of HVT DNA and plasmid DNA into primary chick embryo fibroblast (CEF) cells were done according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUS. The recombinant virus resulting from the cotransfection stock was purified by three successive rounds of plaque purification using the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. When 100% of the plaques were blue, the DNA was analyzed for the presence of the IBDV gene by the SOUTHERN BLOTTING OF DNA procedure. Southern blots, probing EcoRI digested S-HVT-003 DNA with an IBDV specific nick translated probe (plasmid 2-84/2-40), confirmed the presence of the 9100 base pair EcoRI fragment. This result confirmed that S-HVT-003 contained both the lacZ gene and the IBDV gene incorporated into its genome. Additional Southern blots, using a probe specific for BamHI #16, confirmed that the homologous recombination occurred at the appropriate position in BamHI #16 and that no deletions were created. No differences in the growth of S-HVT-003 compared to wild type virus (S-HVT-000) were observed in vitro.

25 Expression of IBDV specific proteins from S-HVT-003 were assayed in vitro using the WESTERN BLOTTING PROCEDURE. Cellular lysates were prepared as described in PREPARATION OF HERPESVIRUS CELL LYSATES. the proteins contained in the cellular lysates of S-30 HVT-003 were separated by polyacrylamide electrophoresis, transferred to introcellulose, and probed with either an antiserum made against denatured purified IBDV capsid proteins or antiserum made against a synthetic peptide corresponding to a predicted imuno 35 dominant region of the IBDV 40 kd (VP2) capsid protein. The filters were washed and treated with [125I] protein A to detect the position of the bound antibodies.

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Figure 4 shows the results obtained using the antiserum made against denatured purified IBDV capsid proteins, which have been shown by the applicants to react primarily with VP3 (32 kd protein). As seen, S-HVT-003 produces protein which a is immunologically indistinguishable from the authentic VP3 protein from intact IBDV virions. Moreover, the polyprotein appears to be processed correctly, producing a VP3 species that comigrates with the authentic VP3 protein. evidence using an Australian IBDV stain indicates that VP4 is involved in the processing of the precursor polyprotein into mature VP2 and VP3 protein species (Jagadish, et al., 1988). Figure 5 shows the results obtained using a rabbit antiserum raised against a synthetic peptide that is homologous to a 14 amino acid region of the IBDV VP2 (40 kd) capsid protein. seen. S-HVT-003 produces a protein is immunologically indistinguishable from the authentic viral VP2 protein. In addition, the VP2 protein produced from S-HVT-003 comigrates with the 40 kd species of VP2 isolated from intact IBDV virions. This species represents a major component of infectious (complete) viral particles.

In summary, analysis of the expression of IBDV specific proteins from S-HVT-003 has shown that the polyprotein is processed in CEF cell culture, producing proteins of the appropriate size that react to immunological reagents specific for either VP2 or VP3 proteins on Western blots.

The following set of experiments was carried out in chickens to analyze the *in vivo* expression of the IBDV genes contained within S-HVT-003 as determined by seroconversion data, serum neutralization results, and protection from IBDV challenge.

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The first experiment was designed to show the seroconversion of chickens to IBDV upon being vaccinated with S-HVT-003. Eleven 11-week-old chickens, seronegative to HVT and IBDV were obtained from SPAFAS Inc. Six birds were vaccinated subcutaneously in the abdominal region with 0.5 ml of a cellular suspension of CEF cells containing S-HVT-003 (40,000 PFU/ml). Serum samples were obtained every seven days for eight weeks for all birds in this study. On day 28 (4th week), three of these birds received a boost of S-HVT-003, while the other three birds received 0.5 ml of an inactivated IBDV vaccine inoculated subcutaneously in the cervical region. Three additional birds were given only the inactivated vaccine on day 28. Two birds served as contact controls and received no vaccinations. On day 56, all birds were sacrificed and necropsied. Table 1 show the results of the serum neutralization assay against IBDV. No detectable SN activity was observed in the birds given only S-HVT-003. Additionally, only one of the three birds that were given only the inactivated vaccine demonstrated low but detectable SN activity. SN titers were also detected in one of the three birds that received the S-HVT-003 followed by the inactivated IBDV vaccine boost; these titers were at a much higher level than with the inactivated IBDV vaccine alone. These results suggest that S-HVT-003 is priming the chicken for a secondary response against IBDV. vitro analysis of the serum samples by WESTERN BLOTTING confirmed the seroconversion of the chickens to IBDV upon vaccination with S-HVT-003 both prior to and after boosts administered on day 28.

DAY

TABLE 1

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5	Vaccine Group	Bird No.	<u>28</u>	31	<u>35</u>	<u>38</u>	42	49
10	HVT-003 HVT-003	266	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 <2 <2
15	HVT-003 IBDV	264	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 1:64 <2	<2 1:256 <2	<2 1:512 <2
20	IBDV•	262	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 <2 <2	<2 1:4 <2	<2 1:4 <2
	С.		<2 <2	<2 <2	<2 <2	<2 <2	<2 <2	<2 <2

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In the second experiment, twenty five 1-day old SPF chicks were vaccinated with S-HVT-003 (20 with 0.2ml 30 subcutaneously and 5 by bilateral eyedrop). chicks were kept as controls. On days four and seven postinfection, five vaccinates and two control birds were bled, sacrificed and their spleens removed for virus isolation. Spleen cell suspensions were made 35 by standard method, and  $\sim 1 \times 10^6$  cells in 3 ml of chick embryo fibroblast (CEF) growth media were inoculated directly onto secondary cells. were incubated for 6-7 days and then scored for cytopathic effects (CPE) as determined by observing cell morphology. The cultures were passed a second time, and again scored for CPE. The results are shown in Table 2. All nonvaccinated control birds remained negative for HVT for both day 4 and 7 spleen cell isolations. Four out of the five birds vaccinated with S-HVT-003 were positive for HVT at

45 day 4 for both the first and second passages. One

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bird did not produce virus, this may represent a vaccination failure. Five out of five birds were positive for HVT on day 7 at both passage one and two. Overall, the vector recovery experiment demonstrates that S-HVT-003 replicates as well as wild type HVT virus in vivo and that insertion of the IBDV/lacZ cassette into the XhoI site of BamHI #16 does not result in detectable attenuation of virus. Subsequent experiments examining the recovered virus by the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure confirmed the in vivo stability of S-HVT-003, by demonstrating  $\beta$ -galactosidase expression in 100% of the viruses.

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TABLE 2
Harvest Date

		<u>Day</u>	<u>74</u>	<u>Da</u>	<u>y 7</u>
	<u>Sample</u>	<u>P1</u>	<u>P2</u>	<u>P1</u>	<u>P2</u>
5	N 1	-	-		
	N 2	-	-		
	N 3			-	-
	N 4			-	-
10	T 1	-	-		
	T 2	2+	2+		
	T 3	2+	2+		
	T 4	+	4+		
	<b>T</b> 5	3+	3+		
15	T 6			2+ cont	taminated
	T 7			+	5+
	T 8			+	5+
	T 8			+	5+
	T 9			+	5+
20	T10			+	5+

N = control, T = vaccinated
CPE ranged from negative (-) to 5+

25 At days 0, 4, 7, 14, 21, and 27 postinfection, blood samples were obtained from the rest of the chickens for determining serum ELISA titers against IBDV and HVT antigens as well as for virus neutralizing tests against IBDV. Additionally, at 21 days postinfection five control and fourteen vaccinated chicks were 30 challenged with virulent IBDV by bi-lateral eyedrop  $(10^{3.8}EID_{50})$ . All birds were sacrificed 6-days post challenge and bursa to body weight ratios A summary of the results is shown in calculated. tables 3 and 4, respectively. As presented in Table 3, 35 no antibodies were detected against HVT antigens by ELISA prior to 21-27 days post vaccination. chickens, the immune response during the first two weeks post hatch is both immature and parentally 40 suppressed, and therefore these results are not totally unexpected. In contrast, IBDV ELISA's were negative up to day 21 post-vaccination, and were only detectable after challenge on day 27. The ELISA levels seen on

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day 27 post-vaccination indicate a primary response to IBDV. Table 4 comparing the Bursa-to-Body weight ratios for challenged controls and vaccinated/challenged groups show no significant differences. Vaccination with S-HVT-003 under these conditions did not prevent infection of the vaccinated birds by IBDV challenge, as indicated by the death of four vaccinated birds following challenge.

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TABLE 3

			1	<u>ELISA</u>	<u>VN</u>
	Sample	Group	HVT	<u>IBDV</u>	<u>IBDV</u>
	C-0	(n=3)	0	0	<100
5	C-4	(n=2)	0	0	nd
	T-4	(n=5)	0	0	nd
	C-7	(n=2)	0	0	<100
	T-7	(n=5)	0	0	<100
	C-14	(n=5)	0	0	nd
10	T-14	(n=14)	0	0	<100
	C-21	(n=5)	0	0	nd
	T-21	(n=14)	1	0	<100
	C-27	(n=5)	0	0	nd
	CC-27	(n=5)	0	5	nd
15	CT-27	(n-10)	3.2	2	nd

C=control

T=vaccinated

CC=challenged control

20 CT=Challenged & vaccinated.

ELISA titers are GMTs and they range from 0-9.

TABLE 4

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23	Sample Group	Body wt.	Bursa wt.	BBR
30	Control (n=5) Challenge Control (n=5)	258.8 209	1.5088 0.6502	0.0058 0.0031
	Challenge Treated (n=10)	215.5	0.5944	0.0027

Values are mean values. Body weights are different in control group because challenged birds did not feed well. Four challenged-treated birds died.

A third experiment was conducted repeating Experiment

2 but using immunologically responsive chicks (3 weeks
of age). Six three week old SPF leghorn chickens were
vaccinated intraperitoneally with 0.2ml of S-HVT-003

(one drop in each eye). Serum samples were obtained
every seven days for six-weeks and the birds were

challenged with the virulent USDA standard challenge

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IBDV virus on day 43 post-vaccination. Six days post challenge, the control, vaccinated-challenged, challenged groups were sacrificed and bursas were harvested for probing with anti-IBDV monoclonal antibodies (MAB) (provided by Dr. David Snyder, Virginia-Maryland Regional College of Veterinary Medicine). Bursal homogenates were prepared by mixing 1 ml of 0.5% NP40 with one bursa. Bursa were then ground and briefly sonicated. Supernatants from the homogenates were reacted with the R63 MAB which had been affixed to 96-well Elisa plates via a protein A linkage. After incubation. a biotin labeled preparation of the R63 MAB was added. After washing, an avidin-horse radish peroxidase conjugate was added and incubated. Tests were developed with Tris-malcate buffer (TMB) +  $H_2O_2$  substrate. The test results are presented in Table 5. The data show the presence of high levels of IBDV antigen in all bursa in the vaccinate-challenged group and in the challenged group. No IBDV antigen was detected in the controls. specific antigen could be detected at dilutions of over 1/1000, and there does not appear to be differences between vaccinated and non-vaccinated challenged HVT titers as determined by ELISA were first detectable at day 7 in four out of the six birds vaccinated. By day 14, six out of six vaccinated birds showed titers to HVT. All six birds continued to show HVT titers throughout the experiment. No IBDV SN titers were seen prior to the challenge. In contrast, analysis of these same serum samples by the WESTERN BLOTTING procedure demonstrated the seroconversion of chickens vaccinated with S-HVT-003 to IBDV prior to administration of the virus challenge. The level of response, however, remains small unless boosted by challenge. Comparison between the vaccinated/challenged and challenged only clearly demonstrates that the level of reactivity by

Western blots is much higher in the vaccinated/challenged group. These results show that S-HVT-003 is seroconverting vaccinated birds to IBDV, and suggest that the level of IBDV specific expression are not high enough to induce a neutralizing response in the birds.

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S-HVT-003 shows the merit of the vaccine approach the applicants have invented. HVT has been engineered to simultaneously express the foreign antigens ( $\beta$ -galactosidase and IBDV antigens) that are recognized in the host by an immune response directed to these proteins.

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TABLE	5
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Serology: Herpes/IBDV ELISA titer

# Bleed Date

5 Bird# 11/3 11/10 11/14 11/24 12/1 12/8 12/15 12/22

# Vaccinated and Challenged

	221	0/0	7/0	5/0	6/0	5/0	5/0	5/0	3/3
	41	0/0	4/0	4/0	1/0	1/0	1/0	1/0	1/3
10	42	0/0	3/0	2/0	1/0	5/0	5/0	5/0	3/2
	43	0/0	0/0	5/0	5/0	5/0	5/0	3/0	3/2
	44	0/0	1/0	5/0	1/0	2/0	1/0	1/0	2/4
	45	0/0	0/0	1/0	1/0	1/0	1/0	1/0	1/3

# Control

15	28	0/0		0/0
	38	0/0		0/0
	73	0/0	·	0/0
	75	0/0		0/0

# Challenged only

20	40	0/0		0/3
	74	0/0		0/5
	39	0/0		0/3
	72	0/0	•	0/3

Maximum titer level is 9

# Example 3

# S-HVT-004

S-HVT-004 is a recombinant herpesvirus of turkeys that contains the Marek's disease virus (MDV) glycoprotein A (gA) gene inserted into the long unique region, and the  $\beta$ -galactosidase (lacZ) gene also inserted in the long unique region. The MDV antigen is more likely to elicit the proper antigentic response than the HVT equivalent antigen.

The MDV gA (SEQ ID NOS: 8 and 9) gene was cloned by standard DNA cloning gA procedures. An EcoRI restriction fragment had been reported to contain the MDV gA gene (Isfort et al., 1984) and this fragment was identified by size in the DNA clones. The region of the DNA reported to contain the gA gene was sequenced by applicants and found to contain a glycoprotein gene as expected. The DNA from this gene was used to find the corresponding gene in HVT by the SOUTHERN BLOTTING OF DNA procedure, and a gene in HVT was identified that contained a very similar sequence. This gene is the same gene previously called gA (Isfort et al., 1984).

For insertion into the genome of HVT, the MDV gA gene was used intact because it would have good herpesvirus signal sequences already. The lacZ gene was inserted into the XhoI fragment in BamHI fragment #16, and the MDV gA gene was inserted behind lacZ as shown in Figures 6A and 6B. Flanking regions in BamHI #16 were used for the homologous recombination. HVT DNA and plasmid DNA were co-transfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUS procedure into primary chick embryo fibroblast (CEF) cells. The virus from the transfection stock was purified by successive plaque purifications using the

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BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. At the end of this procedure, when 100% of the plaques were blue, the DNA was analyzed for the presence of the MDV gA gene. S-HVT-004 is a recombinant virus that contains both the  $\beta$ -galactosidase gene and the MDV gA gene incorporated into the genome.

Figure 6C shows the structure of S-HVT-004.

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# Example 4

# NEWCASTLE DISEASE VIRUS

Newcastle disease virus (NDV) is closely related to PI-3 in overall structure. Hemagglutinin (HN) and fusion (F) genes of PI-3 was engineered for expression in IBR (ref). Similarly hemagglutinin (HN) and fusion (F) genes was cloned from NDV for use in the herpesvirus delivery system (Herpesvirus of turkeys, HVT).

The procedures that was utilized for construction of herpesvirus control sequences for expression have been applied to NDV.

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#### INFECTIOUS BRONCHITIS VIRUS

Infectious bronchitis virus (IBV) is a virus of chickens closely related in overall structure to TGE.

Major neutralizing antigen of TGE was engineered for expression in PRV (ref). Similarly major neutralizing antigens was cloned from three strains of IBV:

Massachusetts (SEQ ID NOs: 14 and 15), Connecticut (SEQ ID NOs: 18 and 19), and Arkansas-99 (SEQ ID NOs: 16 and 17) for use in a herpesvirus delivery system (HVT).

The procedures that was utilized for the construction of herpesvirus control sequences for expression have been applied to IBV.

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#### EXAMPLE 5

# S-HVT-045

S-HVT-045 is a recombinant herpesvirus of turkeys that contains the Marek's disease virus (MDV) glycoprotein B (gB) gene inserted into the short unique region. The MDV antigen is more likely to elicit the proper antigenic response than the HVT equivalent antigen. S-HVT-045 has been deposited on October 15, 1992 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR 2383.

The MDV gB gene was cloned by standard DNA cloning procedures. The MDV gB gene was localized to a 3.9 kb EcoRI-SalI fragment using an oligonucleotide probe based on the HSV gB sequence in a region found to be conserved among known herpesvirus gB genes. The restriction map 3.9 kb EcoRI-SalI fragment is similar to the published map (Ross et al., 1989).

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For insertion into the HVT genome, the MDV gB was used intact because it would have good herpesvirus signal sequences already. The MDV gB gene was inserted into a cloned 17.15 kb BamHI-EcoRI fragment derived from the HVT BamHI #1 fragment. The site used for insertion was the StuI site within HVT US2, previously utilized for the construction of S-HVT-012. The site was initially altered by insertion of a unique HindIII linker, and the MDV gB gene was inserted by standard DNA cloning procedures. Flanking regions in the 17.15 kb BamHI-EcoRI fragment were used, together with the remaining cloned HVT fragments using the PROCEDURE FOR GENERATING

RECOMBINANT HERPESVIRUSES FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The virus obtained from the transfection stock was plaque purified and the DNA was analyzed for the presence of the MDV gB gene. S-HVT-045 is a recombinant virus that contains the MDV gB gene incorporated into the genome at the StuI site in HVT US2 gene.

#### TESTING OF RECOMBINANT S-HVT-045

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studies Two were conducted to demonstrate the effectiveness of these recombinant HVT/MDV viruses in protecting against challenge with virulent Marek's disease virus. In Study A, one-day-old specific pathogen free (SPF) chicks were vaccinated with either S-HVT-045 or S-HVT-046. Seven days post-vaccination, vaccinated chicks, and non-vaccinated, control chicks were challenged with the highly virulent MD-5 strain of Marek's disease virus. Following a 6-week challenge observation period for clinical signs typical of Marek's disease, all chicks were necropsied and examined for lesions diagnostic of Marek's disease. The results, in Table 6, show that both recombinant viruses gave complete protection against a challenge that caused Marek's disease in 90% of non-vaccinated control chicks.

In a second study, one-day-old chicks were vaccinated either with S-HVT-045 or S-HVT-047. A third group of chicks were vaccinated with a USDA-licensed, conventional vaccine comprised of HVT and SB-1 viruses. Five days post-vaccination, the vaccinated chicks and group of non-vaccinated, control chicks were challenged with virulent Marek's virus, strain RB1B. The chicks were observed for 8 weeks for clinical signs of Marek's disease, then necropsied and observed for Marek's lesions. This study demonstrated the ability

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of HVT-045 and HVT-047 to provide 100% protection against challenge (Table 1). The commercial vaccine gave 96% protection, and 79% of the non-vaccinated chicks developed Marek's disease.

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TABLE 6 EFFICACY OF RECOMBINANT HVT/MDV VIRUSES TO PROTECT SUSCEPTIBLE CHICKS AGAINST VIRULENT MAREK'S DISEASE VIRUS

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Marek's Protection

	Vaccine Group	MD-5 Challenge	RB1B Challenge
	S-HVT-045	20/20	24/24
	S-HVT-046	20/20	Not Tested
	S-HVT-047	Not Tested	24/24
15	HVT*	Not Tested	24/25
	Controls	2/20	5/24

a Commercial

# Example 6

# S-HVT-012

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S-HVT-012 is a recombinant herpesvirus of turkeys that contains the  $E.\ coli\ \beta$ -galactosidase (lacZ) gene inserted into the short unique region. The lacZ gene was used to determine the viability of this insertion site in HVT [ATCC F-126 ("Calnek")]. S-HVT-012 has been deposited on October 15, 1992 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure on with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR 2382.

For insertion into the genome of HVT, galactosidase gene was introduced into the unique StuI site of the cloned EcoRI fragment #7 of HVT, i.e., the fragment containing the StuI site within the US2 gene (as described in Methods and Materials). Flanking regions of EcoRI fragment #7 were used for homologous recombination. HVT DNA and plasmid DNA were co-transfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT VIRUS procedure into primary chick embryo fibroblast (CEF) cells. A blue virus obtained from the transfection stock was purified by successive plaque purifications using the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. end of this procedure, when 100% of the plagues were blue, the DNA was analyzed for the presence of the lacZ gene. S-HVT-012 is a recombinant virus that contains the lacZ gene incorporated into the genome at the Stul site within the US2 gene of HVT.

S-HVT-012 may be formulated as a vaccine in the same

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manner as S-HVT-045. When administered to chickens, such a vaccine provides protection against Marek's disease virus.

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### Example 7

# Sites for Insertion of Foreign DNA into HVT

In order to define appropriate insertion sites, a 10 library of HVT BamHI and EcoRI restriction fragments was generated. Several of these restriction fragments (BamHI fragments #16 and #13, and EcoRI fragments #6, and #9 (see figure 1)) were subjected 15 restriction mapping analysis. One unique restriction site was identified in each fragment as a potential insertion site. These sites included XhoI in BamHI fragments #13 and #16, and EcoRI fragment #9 and SalI in EcoRI fragment #6 and StuI in EcoRI fragment #7. A 20  $\beta$ -galactosidase (lacZ) marker gene was inserted in each of the potential sites. A plasmid containing such a foreign DNA insert may be used according to the DNA COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES to CONSTRUCT a HVT containing the foreign DNA. this procedure to be successful it is important that 25 the insertion site be in a region non-essential to the replication of the HVT and that the site be flanked with HVT DNA appropriate for mediating homologous recombination between virus and plasmid DNAs. 30 plasmids containing the lacZ marker gene were utilized in the DNA COTRANSFECTION FOR GENERATING RECOMBINANT HERPESVIRUSES. The generation of recombinant virus was determined by the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS. Three of the five sites were successfully 35 used to generate a recombinant virus. In each case the resulting virus was easily purified to 100%, clearly defining an appropriate site for the insertion of

foreign DNA. The three homology vectors used to define these sites are described below.

# Example 7A

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# Homology Vector 172-29.31

The homology vector 172-29.31 contains the HVT BamHI #16 fragment and is useful for the insertion of foreign DNA into HVT. Plasmid 172-29.31 contains a unique XhoI restriction site into which foreign DNA may be cloned. XhoI site in homology vector 172-29.31 may be used to insert foreign DNA into HVT by the construction of at least three recombinant HVT (see examples 1-3).

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The homology vector 172-29.31 was further characterized by DNA sequence analysis. The complete sequences of the BamHI #16 fragment was determined. Approximately 2092 base pairs of the adjacent BamHI #13 fragment was also determined (see SEQ ID NO: 3). This sequence indicates that the open reading frame coding for HVT glycoprotein A (gA) spans the BamHI #16 - BamHI #13 junction. The HVT gA gene is homologous to the HSV-1 glycoprotein C (gC). The XhoI site interrupts an ORF which lies directly upstream of the HVT gA gene. This ORF shows amino acid sequence homology to the PRV p43 and the VZV gene 15. The PRV and VZV genes are the homologues of HSV-1 UL43. Therefore this ORF was designated as HVT UL43 (SEQ ID NO: 5). It should be noted that the HVT UL43 does not exhibit direct homology to HSV-1 UL43. Although HVT UL43 is located upstream of the HVT gC homologue it is encoded on the same DNA strand as HVT gA, where as the HSV-1 UL43 is on the opposite strand relative to HSV-1 gC. The XhoI site interrupts UL43 at approximately amino acid 6, suggesting that the UL43 gene is non-essential for HVT replication.

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# Example 7B

# Homology Vector 435-47.R17

The homology vector 435-47.R17 contains the HVT EcoRI #7 fragment and is useful for the insertion of foreign DNA into HVT. Plasmid 435-47.R17 contains a unique HindIII restriction site into which foreign DNA may be cloned. The HindIII restriction site in plasmid results from the insertion of a HindIII linker into the naturally occurring StuI site of EcoRI fragment #7. HindIII site in homology vector 435-47.R17 may be used to insert foreign DNA into HVT by the construction of at least 25 recombinant HVT.

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DNA sequence analysis at the StuI indicated that this fragment contains open reading frames coding for US10, US2, and US3. The StuI site interrupts US2 at approximately amino acid 124, suggesting that the US2 gene is non-essential for HVT replication.

# Example 7C

# Homology Vector 172-63.1

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The homology vector 172-63.1 contains the HVT EcoRI #9 fragment and is useful for the insertion of foreign DNA into HVT. Plasmid 172-63.1 contains a unique XhoI restriction site into which foreign DNA may be cloned. XhoI site in homology vector 172-63.1 may be used to insert foreign DNA into HVT by the construction of S-HVT-014 (see example 8).

# Example 8

# S-HVT-014

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S-HVT-014 is a recombinant herpesvirus of turkeys that contains the  $E.~coli~\beta$ -galactosidase (lacZ) gene inserted into the long unique region. The lacZ gene was used to determine the viability of this insertion site in HVT [ATCC F-126 ("Calnek")].

insertion into the genome of HVT, the galactosidase gene was introduced into the unique XhoI site of the cloned EcoRI fragment #9 (as described in Methods and Materials). The XhoI site within the EcoRI #9 fragment of the HVT genome is the same site as the XhoI site within the BamHI #10 fragment used for construction recombinant herpesvirues of turkevs described in Examples 16 through 19. Flanking regions ECORI fragment #9 were used for homologous recombination. HVT DNA and plasmid DNA were cotransfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT VIRUS procedure into primary chick embryo fibroblast (CEF) cells. A blue virus obtained from the transfection stock was purified by successive plaque purifications using the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. At the end of this procedure when 100% of the plaques were blue. S-HVT-014 is a recombinant virus that contains the lacZ gene incorporated into the genome at the XhoI site within the EcoRI #9 fragment of HVT.

S-HVT-014 may be formulated as a vaccine in the same manner as S-HVT-045. When administered to chickens, such a vaccine provides protection against Marek's disease virus.

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## Example 9

#### S-HVT-005

S-HVT-005 is a recombinant herpesvirus of turkeys that contains the  $E.~coli~\beta$ -galactosidase (lacZ) gene inserted into the long unique region. The lacZ gene was used to determine the viability of this insertion site in HVT [ATCC F-126 ("Calnek")].

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For insertion into the genome of HVT. the galactosidase gene was introduced into an approximately 1300 base pair deletion of the XhoI #9 fragment of HVT. The deletion which lies between the unique MluI and EcoRV sites removes the complete coding region of the HVT gA gene (see SEQ ID NO: 3). Flanking regions of XhoI fragment #9 were used for homologous recombination. HVT DNA and plasmid DNA were cotransfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT VIRUS procedure into primary chick embryo fibroblast (CEF) cells. A blue virus obtained from the transfection stock was purified by successive plaque purifications using the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. At the end of this procedure, when 100% of the plagues were blue, the DNA was analyzed for the presence of the lacZ gene. S-HVT-005 is a recombinant virus that contains the lacZ gene incorporated into the genome in place of the deleted gA gene of HVT.

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S-HVT-005 may be formulated as a vaccine in the same manner as S-HVT-045. When administered to chickens, such a vaccine provides protection against Marek's disease virus.

## Example 10

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## Marek's Disease Vaccines

Recombinant HVT expressing glycoproteins from Marek's Disease Virus make superior vaccines for Marek's Disease. We have constructed several recombinant HVT expressing MDV glycoproteins: S-HVT-004 (Example 3), S-HVT-045 (Example 5), S-HVT-046 (Example 10A), S-HVT-047 (Example 10B), S-HVT-062 (Example 10C).

## Example 10A S-HVT-046

S-HVT-046 is a recombinant herpesvirus of turkeys that contains the Marek's disease virus (MDV) glycoprotein B (gB) and glycoprotein A (gA) genes inserted into the short unique region. The MDV genes are inserted in the same transcriptional orientation as the US2 gene. The MDV antigens are more likely to elicit the proper antigenic response than the HVT equivalent antigen.

S-HVT-046 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 456-17.22 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

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## Example 10B S-HVT-047

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S-HVT-047 is a recombinant herpesvirus of turkeys that contains the MDV gB and gA genes inserted into the short unique region. The MDV genes are inserted in the opposite transcriptional orientation as the US2 gene. The MDV antigens are more likely to elicit the proper antigenic response than the HVT equivalent antigen.

S-HVT-047 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 456-17.18 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

## 20 <u>Example 10C</u> <u>S-HVT-062</u>

S-HVT-062 is a recombinant herpesvirus of turkeys that contains the MDV gB, glycoprotein D (gD) and gA genes inserted into the short unique region. The MDV genes are inserted in the same transcriptional orientation as the US2 gene. The MDV antigens are more likely to elicit the proper antigenic response than the HVT equivalent antigen. S-HVT-062 has been deposited on February 23, 1993 pursuant to the Budapest Treaty on the International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR 2401.

S-HVT-062 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC

DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 556-60.6 with BamHI and HindIII, and 456-17.22 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

## TESTING OF RECOMBINANT HVT EXPRESSING MDV ANTIGENS

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Two studies were conducted to demonstrate the effectiveness of these recombinant HVT/MDV viruses in protecting against challenge with virulent Marek's disease virus. In Study 1, one-day-old specific pathogen free (SPF) chicks were vaccinated with either S-HVT-045, S-HVT-046, or S-HVT-047. Five days postvaccination, vaccinated chicks, and non-vaccinated, control chicks were challenged with MDV. Following a 6week post-challenge observation period for clinical signs typical of Marek's disease, all chicks were necropsied and examined for lesions diagnostic of Marek's disease. The results, in Table 7, show these recombinant viruses gave complete protection against a challenge that caused Marek's disease in 84% of nonvaccinated control chicks.

In the second study, one-day-old chicks were vaccinated with S-HVT-062. Two more groups of chicks were vaccinated with a USDA-licensed, conventional vaccines comprised of HVT and a combination HVT and SB-1 viruses. Five days post-vaccination, the vaccinated chicks and a group of non-vaccinated, control chicks were challenged with MDV. The chicks were observed for

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8 weeks for clinical signs of Marek's disease, then necropsied and observed for Marek's lesions. This study demonstrated the ability of S-HVT-062 to provide 100% protection against challenge (Table 7). The commercial vaccines gave 81% and 95% protection, respectively and 100% of the non-vaccinated chicks developed Marek's disease.

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# TABLE 7 EFFICACY OF RECOMBINANT HVT/MDV VIRUSES AGAINST VIRULENT MAREK'S VIRUS CHALLENGE

5	Study	Vaccine Group	Dose*	Protection <sup>b</sup>
	1	S-HVT-045	2.2 X 10 <sup>3</sup>	24/24 (100%)
	1	S-HVT-046	2.2 X 10 <sup>3</sup>	20/20 (100%)
10	1	S-HVT-047	2.2 X 10 <sup>3</sup>	24/24 (100%)
	1	Controls		7/44 (16%)
15	1	HVT/SB-1		24/25 (96%)
	2	S-HVT-062	7.5 X 10 <sup>2</sup>	32/32 (100%)
	2	S-HVT-062	1.5 X 103	22/22 (100%)
20	2	Controls		0/20 (0%)
	2	HVT°	7.5 X 10 <sup>2</sup>	17/21 (81%)
25	2	HVT/SB-1°	7.5 X 10 <sup>2</sup>	21/22 (95%)

<sup>•</sup> PFU/0.2 ml.

No. protected/Total; Challenge 5 days postvaccination.

<sup>30 °</sup> Commercial vaccine.

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## Example 11

## Bivalent Vaccines Against Newcastle Disease and Marek's Disease

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Recombinant HVT expressing proteins from NDV make bivalent vaccines protecting against both Marek's Disease and Newcastle disease. Several recombinant HVT expressing NDV proteins were constructed S-HVT-007 (Example 11A), S-HVT-048 (Example 11B), S-HVT-049 (Example 11C), S-HVT-050 (Example 11D), and S-HVT-106 (Example 11E).

## Example 11A S-HVT-007

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S-HVT-007 is a recombinant herpesvirus of turkeys that contains a  $E.\ coli$  lacZ NDV HN hybrid protein gene under the control of the PRV gX promoter and the NDV F gene under the control of the HSV-1  $\alpha 4$  promoter inserted into the long unique region. The NDV genes are inserted in the same transcriptional orientation as the UL43 gene.

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To construct S-HVT-007, HVT DNA and the plasmid 255-18.B16 were co-transfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT VIRUS procedure into primary chick embryo fibroblast (CEF) cells. A blue virus obtained from the transfection stock was purified by successive plaque purifications using the BLUOGAL SCREEN FOR RECOMBINANT HERPESVIRUS procedure. At the end of this procedure, when 100% of the plaques were blue.

## Example 11B S-HVT-048

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S-HVT-048 is a recombinant herpesvirus of turkeys that contains the MDV gB and gA genes and the NDV F gene under the control of the HCMV immediate early promoter inserted into the short unique region. The MDV and NDV genes are inserted in the same transcriptional orientation as the US2 gene.

S-HVT-048 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 535-70.3 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

## 20 Example 11C S-HVT-049

S-HVT-049 is a recombinant herpesvirus of turkeys that contains the MDV gB and gA genes and the NDV HN gene under the control of the PRV gX promoter inserted into the short unique region. The MDV and NDV genes are inserted in the same transcriptional orientation as the US2 gene.

S-HVT-049 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 549-62.10 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

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## Example 11D S-HVT-050

S-HVT-050 is a recombinant herpesvirus of turkeys that contains the MDV gB and gA genes and the NDV HN (SEQ ID NOs: 10 and 11) and F (SEQ ID NOs: 12 and 13) genes. The NDV genes are under the control of the PRV gX and HCMV immediately promoters respectively. All four genes are inserted into the short unique region in the same transcriptional orientation as the US2 gene.

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S-HVT-050 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 549-24.15 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis. S-HVT-050 has been deposited on February 23, 1993 pursuant to the Budapest Treaty International Deposit of Microorganisms for the Purposes of Patent Procedure with the Patent Culture Depository of the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 U.S.A. under ATCC Accession No. VR 2400.

#### Example 11E S-HVT-106

S-HVT-106 is a recombinant herpesvirus of turkeys that contains the MDV gA, gB, gD genes and the NDV HN and F genes. The NDV genes are under the control of the PRV gX and HCMV immediately promoters respectively. All five genes are inserted into the short unique region in the same transcriptional orientation as the US2 gene.

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S-HVT-106 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC

DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 633-13.27 uncut.

#### TESTING OF RECOMBINANT HVT EXPRESSING NDV ANTIGENS

studies were conducted to demonstrate effectiveness of these recombinant HVT/MDV/NDV viruses in protecting against challenge with virulent Newcastle and Marek's disease viruses. In Study 1, one-day-old specific pathogen free (SPF) chicks were vaccinated with either S-HVT-048. S-HVT-049, S-HVT-050, or a USDA-licensed, conventional vaccine comprised of NDV B1/B1 virus. Three weeks post-vaccination, vaccinated chicks. and non-vaccinated, control chicks challenged with NDV. Birds were then observed for clinical signs of disease. The results, in Table 8, show these recombinant viruses (S-HVT-048 and S-HVT-050) gave complete protection against a challenge that caused Newcastle disease in 100% of non-vaccinated Recombinant virus S-HVT-049 control chicks. partial protection against Newcastle disease.

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In the second study, one-day-old chicks were vaccinated with S-HVT-050. Two more groups of chicks were vaccinated with a USDA-licensed, conventional vaccines comprised of HVT and a combination HVT and SB-1 viruses. Five days post-vaccination, the vaccinated chicks and a group of non-vaccinated, control chicks were challenged with MDV. The chicks were observed for 8 weeks for clinical signs of Marek's disease, then necropsied and observed for Marek's lesions. This study demonstrated the ability of S-HVT-050 to provide protection greater than the commercial Marek's disease vaccines.

TABLE 8 EFFICACY OF RECOMBINANT HVT/MDV/NDV VIRUSES AGAINST VIRULENT NEWCASTLE AND MAREK'S DISEASE VIRUS CHALLENGE

Protection (%)

5			11000011011 (0)			
	Study	Vaccine Group	Dose*	NDV <sup>b</sup>	MDV°	
10	1	S-HVT-048	4.0 X 10 <sup>4</sup>	19/19 (10	0)	
	1	S-HVT-049	3.0 X 104	4/20 (20)		
15	1	S-HVT-050	1.5 X 104	20/20 (10	0)	
13	1	Controls		0/20 (0)		
	1	NDV B1/B1d		18/18 (10	0)	
20	2	S-HVT-050	$7.5 \times 10^{2}$		13/14 (93)	
	2	S-HVT-050	$1.5 \times 10^3$		16/17 (94)	
25	2 .	Controls			5/23 (22)	
23	2 H	HVTd			20/26 (77)	
	2	HVT/SB-1d			10/12 (83)	
30	a PFU/	0.2 ml.				
	b No. p	rotected/Total	; Challenge 3	weeks post-v	accination.	
<b>3</b> E	c No. p	rotected/Total	; Challenge 5	days post-va	ccination.	
35	d Comme	rcial vaccine.				

## Example 12

## Bivalent Vaccines Against Infectious Laryngotracheitis and Marek's Disease

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Recombinant HVT expressing glycoproteins from ILT virus make bivalent vaccines protecting against both Marek's disease and infectious laryngotracheitis. Several recombinant HVT expressing ILT virus glycoproteins S-HVT-051 (Example 12A), S-HVT-052 (Example 12B), and S-HVT-104 (Example 11C) were constructed.

## Example 12A S-HVT-051

- S-HVT-051 is a recombinant herpesvirus of turkeys that contains the ILT virus gB gene inserted into the short unique region. The ILT gene is inserted in the same transcriptional orientation as the US2 gene.
- S-HVT-051 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 528-11.34 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

## 30 **Example 12B** S-HVT-052

S-HVT-052 is a recombinant herpesvirus of turkeys that contains the ILT virus gD gene inserted into the short unique region. The ILT gene is inserted in the opposite transcriptional orientation as the US2 gene.

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S-HVT-052 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 437-26.26 with BamHI and HindIII, and 528-03.37 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

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## Example 12C S-HVT-104

S-HVT-104 is a recombinant herpesvirus of turkeys that contains six foreign genes. The MDV gA, gB, and gD genes are inserted in the unique short region in the same transcriptional orientation as the US2 gene. An E. coli lacZ marker gene and the ILT gB and gD genes are inserted in BamHI #16 region in the same transcriptional orientation as the UL43 gene.

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To construct S-HVT-104, DNA from S-HVT-062 and the plasmid 634-29.16 were co-transfected according to the DNA TRANSFECTION FOR GENERATING RECOMBINANT VIRUS procedure into primary chick embryo fibroblast (CEF) cells.

TESTING OF RECOMBINANT HVT EXPRESSING ILT ANTIGENS

The following study was conducted to demonstrate the effectiveness of these recombinant HVT/ILT viruses in protecting against challenge with virulent Infectious Laryngotracheitis virus. One-day-old specific pathogen free (SPF) chicks were vaccinated with either S-HVT-051, S-HVT-052, a combination of S-HVT-051 and S-HVT-052, or a USDA-licensed, conventional vaccine comprised of ILT virus. Two to three weeks post-vaccination, vaccinated chicks, and non-vaccinated, control chicks

were challenged with ILT. Birds were then observed for clinical signs of disease. The results, in Table 9, show these recombinant viruses (S-HVT-051 and S-HVT-052) gave protection against challenge with ILT virus comparable to a commercial ILT vaccine.

Animals vaccinated with the vaccines described here may be easily differentiated from animals infected with virulent ILT. This is accomplished by testing the suspect birds for antibodies to any ILT antigens other than gB or gD. Examples of such antigens are ILT glycoproteins C, E, and G. Vaccinated, uninfected birds will be negative for these antigens whereas infected birds will be positive.

TABLE 9 EFFICACY OF RECOMBINANT HVT/ILT VIRUSES AGAINST VIRULENT INFECTIOUS LARYNGOTRACHEITIS VIRUS CHALLENGE

5	Vaccine Group	Dose*	Protection <sup>b</sup>
	S-HVT-051		28/30 (93%)
		2.1 X 10 <sup>3</sup>	
	S-HVT-052	1.7 X 10 <sup>3</sup>	29/29 (100%)
	S-HVT-051 +	2.1 X 103	24/24 (100%)
	S-HVT-052	1.7 X 10 <sup>3</sup>	
10	Controls	2/30 (7%)	
	ILT <sup>c</sup>		29/30 (97%)

PFU/0.2 ml.

No.protected/Total; Challenge 2-3 weeks postvaccination.

c Commercial vaccine.

## Example 13

## Bivalent Vaccines Against Infectious Bursal Disease and Marek's Disease

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Recombinant HVT expressing proteins from IBDV make bivalent vaccines protecting against both Marek's and infectious bursal disease. Disease Several expressing recombinant HVT IBDV proteins were These viruses include S-HVT-003 (example constructed. 2) and S-HVT-096.

S-HVT-096 is a recombinant herpesvirus of turkeys that contains the IBDV VP2 gene, under the control of the HCMV immediate early promoter, inserted into the short unique region. The IBDV gene is inserted in the same transcriptional orientation as the US2 gene.

S-HVT-096 was constructed according to the PROCEDURE 20 FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with Not1. 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 556-60.6 25 with BamHI, and 602-57.F1 uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.

S-HVT-096 was assayed for expression of VP2 by black plaque and western blot analysis. Both assays indicated that the virus was expressing high levels of protein which reacts specifically with an neutralizing monoclonal antibody. This virus will be useful as a vaccine against infectious bursal disease.

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## Example 14

# <u>Bivalent Vaccines Against Infectious Bronchitis and Marek's Disease</u>

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S-HVT-066 is a recombinant herpesvirus of turkeys that contains the MDV gB, gD and gA genes and the IBV spike and matrix genes. The IBV spike and matrix genes are under the control of the HCMV immediate early and PRV gX promoters respectively. All five genes are inserted into the short unique region. The MDV and IBV genes are inserted in the same transcriptional orientation as the US2 gene.

- S-HVT-066 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM SUBGENOMIC DNA FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 407-32.1C1 with NotI, 437-26.24 with BamHI and HindIII, 556-60.6 with BamHI, and 567-72.1D uncut. Insertion of the appropriate DNA was confirmed by southern blot analysis.
- S-HVT-066 was assayed for expression of the IBV spike protein by black plaque and western blot analysis. Both assays indicated that the virus was expressing high levels of protein which reacts specifically with an IBV neutralizing monoclonal antibody. This virus will be useful as a vaccine against infectious bronchitis.

## Example 15

<u>Vaccines utilizing HVT to express antiqens from various pathogens</u>.

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Anticipate that antigens from the following pathogens may also be utilized to develop poultry vaccines: Chick anemia virus (agent), Avian encephalomyelitis virus, Avian reovirus, Avian paramyxoviruses, Avian influenza virus, Avian adenovirus, Fowl pox virus, Avian coronavirus, Avian rotavirus, Salmonella spp E. coli, Pasteurella spp, Haemophilus spp, Chlamydia spp, Mycoplasma spp, Campylobacter spp, Bordetella spp, Poultry nematodes, cestodes, trematodes, Poultry mites/lice, Poultry protozoa (Eimeria spp, Histomonas spp, Trichomonas spp).

## Example 16

20 Trivalent vaccines against Infectious Laryngotracheitis, Marek's Disease and Newcastle's and bivalent vaccines against Infectious Laryngotracheitis and Marek's Disease are described. Superior protection against Infectious 25 Laryngotracheitis is achieved with a vaccine combining S-HVT-123 (expressing ILTV gB and gD) with S-HVT-138, -139, or 140 (expressing ILTV gD and gI).

## Example 16A S-HVT-123

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S-HVT-123 is a recombinant herpesvirus of turkeys that contains the ILT virus gB and gD genes inserted into an XhoI site converted to a NotI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figures 13B and 15; SEQ ID NO: 48). S-HVT-123 further contains the MDV gA, gD, and gB genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The

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ILTV genes and the MDV genes each use their own respective promoters. S-HVT-123 is useful as a vaccine in poultry against Infectious Laryngotracheitis and Marek's Disease.

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S-HVT-123 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 721-38.1J uncut, 729-37.1 with AscI.

## Example 16B S-HVT-138

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S-HVT-138 is a recombinant herpesvirus of turkeys that contains the ILT virus gD and gI genes inserted into a unique XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figures 13A and 15). The ILTV gD and gI genes are in the opposite transcriptional orientation to the open reading frame (ORF A) within the Ecor1 #9 (BamHI #10) fragment of the HVT genome (Figure 14; SEQ ID NOS: 48, 50). The ILTV gD and gI genes are expressed as overlapping transcripts from endogenous ILTV promoters, and share their own endogenous polyadenylation signal.

S-HVT-138 is useful as a vaccine in poultry against Infectious Laryngotracheitis and Marek's Disease.

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S-HVT-138 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 711-92.1A uncut, 415-09.BA1 with BamHI.

Sera from S-HVT-138 vaccinated chickens reacts on Western blots with ILTV gI protein indicating that the S-HVT-138 vaccine expressed the ILTV protein and does elicit an immune response in birds. S-HVT-138 vaccinated chickens were protected from challenge by virulent infectious laryngotracheitis virus.

## Example 16C S-HVT-139

S-HVT-139 is a recombinant herpesvirus of turkeys that contains the ILT virus gD and gI genes inserted into a unique XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome. The ILTV gD and gI genes are in the opposite transcriptional orientation to the open reading frame (ORF A) within the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figure 13A and 15; SEQ ID NO: 48, 50). further contains the MDV gA, gD, and gB genes are inserted into the unique StuI site converted into a HindIII site in the HVT US2 gene. The ILTV gD and gI genes are expressed as overlapping transcripts from their won respective endogenous ILTV promoters, and the MDV genes are also expressed from their own endogenous promoters. S-HVT-139 is useful as a vaccine in poultry against Infectious Laryngotracheitis and Marek's Disease.

S-HVT-139 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 711-92.1A uncut, 721-38.1J uncut.

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S-HVT-140 is a recombinant herpesvirus-of turkeys that contains the ILT virus gD and gI genes inserted into a unique XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figures 13A and 15). The ILTV gD and gI genes are in the opposite transcriptional orientation to the open reading frame (ORF A) within the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figure 14; SEQ ID NO: 48, 50). S-HVT-140 further contains the MDV gA, gD, and gB genes and the NDV F and HN genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The ILTV gD and gI genes are expressed as overlapping transcripts from their own respective endogenous ILTV promoters, and the MDV genes are also expressed from their own respective endogenous MDV promoters. The NDV F gene is transcribed from the HCMV immediate early promoter, and the NDV HN gene is transcribed from the PRV gX promoter. S-HVT-140 is useful as a vaccine in poultry against Infectious Laryngotracheitis, Marek's Disease, and Newcastle's Disease.

S-HVT-140 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 711-92.1A uncut, 722-60.E2 uncut.

## 30 **Example 17**

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Trivalent vaccines against Infectious Bursal Disease, Marek's Disease and Newcastle's Disease and bivalent vaccines against Infectious Bursal Disease and Marek's Disease are described.

#### Example 17A HVT-126

S-HVT-126 is a recombinant herpesvirus of turkeys that contains the IBDV VP2 gene inserted into an XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment in the HVT genome (Figures 13A and 15). The IBDV gene is in the same transcriptional orientation as the open reading frame (ORF A) within the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figure 14; SEQ ID NO: 48, 50). The IBDV VP2 gene is expressed from an IBRV VP8 promoter. S-HVT-126 is useful as a vaccine in poultry against Infectious Bursal Disease and Marek's Disease.

S-HVT-126 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 706-57.A3 uncut, 415-09.BA1 with BamHI.

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## Example 17B HVT-137

S-HVT-137 is a recombinant herpesvirus of turkeys that contains the IBDV VP2 gene inserted into a uniqe XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment in the HVT genome (Figures 13A and 15). The is in gene the same transcriptional orientation as the open reading frame (ORF A) within the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figure 14; SEQ ID NO: 48, 50). S-HVT-137 further contains the MDV gA, gD, and gB genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The IBDV VP2 gene is expressed from an IBRV VP8 promoter. The MDV genes are expressed from their own respective endogenous MDV promoters. S-HVTis useful as a vaccine in poultry against Infectious Bursal Disease and Marek's Disease.

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S-HVT-137 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 706-57.A3 uncut, 721-38.1J uncut.

## Example 17C HVT-143

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S-HVT-143 is a recombinant herpesvirus of turkeys that contains the IBDV VP2 gene inserted into a unique XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figures 13 A and 15). The gene is in the same transcriptional orientation as the open reading frame (ORF A) within the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figure 14; SEQ ID NO: 48, 50). S-HVT-143 further contains the MDV gA, gD, and gB genes and the NDV F and HN genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The IBDV VP2 gene is expressed from an IBRV VP8 promoter. The MDV are expressed from their own respective endogenous MDV promoters. The NDV F gene is transcribed from the HCMV immediate early promoter, and the NDV HN gene is transcribed from the PRV gX promoter. S-HVT-143 is useful as a vaccine in poultry against Infectious Bursal Disease, Marek's Disease, and Newcastle's Disease.

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S-HVT-143 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 706-57.A3 uncut, 722-60.E2 uncut.

## Example 18 HVT-128

S-HVT-128 is a recombinant herpesvirus of turkeys that contains the NDV HN and F genes inserted into a unique XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment of the HVT genome (Figures 13A and 15). S-HVT-128 further contains the MDV gA, gD, and gB genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The NDV HN gene is expressed from the PRV gX promoter and the NDV F gene is expressed from the HCMV immediate early promoter. The MDV genes are expressed from the endogenous MDV promoters. S-HVT-128 is useful as a vaccine in poultry against Newcastle's Disease and Marek's Disease.

S-HVT-128 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, and 717-38.12 uncut. To a mixture of these six cosmids was added a limiting dilution of a recombinant HVT virus containing the MDV gA, gD, and gB genes inserted into the unique short region (see HVT-062) and the PRV gX promoter-lacZ gene inserted into an XhoI site converted to a NotI site in the EcoR1 #9 (BamHI #10) fragment within the unique long region of HVT. A recombinant virus S-HVT-128 was selected which was lac Z negative.

## Example 18B HVT-136

S-HVT-136 is a recombinant herpesvirus of turkeys that contains the NDV HN and F genes inserted into an XhoI site converted to a PacI site in the EcoR1 #9 (BamHI #10) fragment within the unique long region of HVT. (Figure 14; SEQ ID NOs: 48 and 50) The NDV HN gene is

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expressed from the PRV gX promoter and the NDV F gene is expressed from the HCMV immediate early promoter. S-HVT-136 is useful as a vaccine in poultry against Newcastle's disease and Marek's disease.

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S-HVT-136 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, and 717-38.12 uncut, and 415-09.BA1 with BamHI.

## Example 19 S-HVT-145

## HVT/MDV recombinant virus vaccine

S-HVT-145 is a recombinant virus vaccine containing MDV and HVT genomic sequences which protects against Marek's disease is produced by combining cosmids of MDV genomic DNA containing genes coding for the relevant protective antigens of virulent MDV serotype 2 and cosmids of HVT genomic DNA according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The resulting virus is a vaccine that ahs the protective immune respnse to virulent MDV serotype 2 and the attenuated growth characteristics of In one embodiment, a chimeric virus vaccine the HVT. containing the MDV genes of the unique short and the HVT genes of the unique long is useful as a vaccine Marek's disease in chickens. protective antigens withinthe unique short (gD, gE, and qI) elicit a protective immune response to MDV, while the virulence elements present in the unique long of MDV (55,56, 57) are replaced by the attenuating uniuge long sequences of HVT. The result is an attenuated

virus vaccine which protects against Marek's disease. Multivalent protection against Marek's infectious laryngotracheitis, infectious vursal disease, Newcastle's dises, or another poultry pathogen is achieved by inserting the ILTV gB, gD, and gI genes, the IBDV VP2 gene, the NDV HN and F genes, or an antigen gene from apoultry pathogen into an XhoI site converted to a PacI site or NotI site in the EcoR1 #9 (BamHI #10) fragment within the uniuge long region of HVT/MDV recombinant virus (Figures 13 and 15).

A cosmid was constructed containing the entir MDV unique short region. MDV genomic DNa contains several Smal sites in the uniuge long and internal and terminal repeats of the virus, but no SmaI sites wihin the unique short of the virus. The entire unique short region of MDV was isolated by a partial restriction digestion of MDV genomic DNa with Smal. A DNA fragment approximately 29,000 to 33,000 base pairs was isolated and cloned into a blunt ended site of the cosmid vector To generate HVY-145, a recombinant HVT/MDV chimeric virus, the cosmid containing the MDV unique short region was combined with cosmids containing the HVT unique long region according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI. 407-32.1C1 with NotI, and 739-27.16 with NotI.

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The resulting virus vaccine provides superior protection against Marek's disease or as a multivalent vaccine against Marek's disease and infectious laryngotracheitis, infectious bursal disease. Newcastle's disease, or another poultry pathogen. This vaccine is superior because expression of MDV genes in the HVT/MDV chimera vaccine is safer and provides

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better protection against Marke's disease than vaccines presently available containing HVT and MDV type 1 (SBor HVT alone. Secondly, one can demonstrate expression of the MDV glycoprotein gens in the absence of the homologous HVT genes for both diagnostic and regulatory purposes. This is useful since antibodies to an MDV glycoprotein will cross react with the homologous HVT glycoprotein. Finally, a recombinant HVT/MDV virus which contains a single copy of each glycoprotein gene is more stable that a recombinant containing two copies of a homologous glycoprotein gene from HVT and MDV which may delete by homologous recombination.

In an alternative embodiment, cosmids containing MDV protective antigen genes from the unique long (MDV gB and gC) are combined with cosmids containing HVT gene sequences from the unique short and the unique long, effectively avoiding the MDV vírulence genes at the unique long/internal repeat junction and the unique long/terminal repeat junction (55, 56, and 57).

SB-1 strain is an MDV serotype 1 with attenuated pathogenicity. Vaccination with a combination of HVT and SB-1 live viruses protects against virulent MDV challenge better than vaccination with either virus In an alternative embodiment of the present invention, recombinant virus vaccine comprises a protective antigen genes of the virulent MDV serotypes 2 combined with the attenuating genes of the nonvirulent MDV serotypes 1 and 3, such as SB-1 and HVT. The genomic DNA corresponding to the unique long region is contributed by the SB-1 serotype. The genomic DNA corresponding to the unique short region is contributed by the HVT serotype. Three major glycoprotein antigens (gB, gA and gD) from the MDV serotype 2 are inserted into the unique short region of the virus.

The recombinant virus is constructed utilizing HVT subgenomic clones 672-01.A40, 672-07.C40 and 721-38.1J to reconstruct the unique short region. Subgenomic clone 721-38.1J contains an insertion of the MDV gB, qA, and qD genes. A large molar excess of these clones is cotransfected with a sub-infectious dose of Sb-1 To determine the appropriate subgenomic DNA. infectious dose, transfection of the SB-1 is titrated down to a dose which no longer yields virus plaques in Such a dose contains sub-genomic cell culture. fragments spanning the unique long region of SB-1 which recombine withthe HVT unique short subgenomic clones. Therefore, a virus resulting from recombination between overlapping homologous regions of the SB-1 and HVT subgenomic fragments is highly favored. Alternatively, SB-1 genomic fragments from the unique long region are subcloned into cosmid vectors. A recombinant virus containing the Sb-1 unique long the HVT unique short with the MDV, qB, qA, and qD genes were produced using the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. This procedure is also used with an HVT subgenomic clone to insert antigen genes from other avian pathogens including but not limited to infectious laryngotracheitis virus, Newcastle's disease virus and infectious bursal disease virus.

### Example 20

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Recombinant HVT expressing chicken myelomonocytic growth factor (cMGF) or chicken interferon (cIFN) are useful as vaccines against Marek's disease virus and are also useful to enhance the immune response against other diseases of poultry. Chicken myelomonocytic growth factor (cMGF) is related to mammalian G-CSF and interleukin-6 protein (58), and chicken interferon (cIFN) is homologous to mammalian type 1 interferon

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(59) interferon. When used in combination with vaccines described in previous examples, S-HVT-144 or HVT expressing cIFN are useful to provide enhanced mucosal, humoral, or cell mediated immunity against avian disease-causing viruses including, but not limited to, Marek's disease virus, Newcastle disease virus, infectious laryngotracheitis virus, infectious bronchitis virus, infectious bursal disease virus. Recombinant HVT expressing cMGF or cIFN are useful provide enhanced immunity against avian disease causing organismsdescribed in Example 15.

## Example 20A S-HVT-144

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S-HVT-144 is a recombinant herpesvirus of turkeys that contains the chicken myelomonocytic growth factor (cMGF) gene inserted into an XhoI site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT. The cMGF gene is in the opposite transcriptional orientation to the open reading frame (ORF A) within the EcoR1 #9 fragment of the HVT genome (Figure 14; SEQ ID NOs: 48 and 50). The cMGF gene is expressed from a human cytomegalovirus immediate early promoter. S-HVT-144 is useful as a vaccine in poultry against Marek's Disease.

S-HVT-144 was constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 751-87.A8 with Asc I, 415-09.BA1 with BamHI.

35 <u>Example 20B</u> Recombinant HVT expressing chicken interferon

A recombinant herpesvirus of turkeys contains the chicken interferon (cIFN) gene inserted into an XhoI site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT. The cIFN gene is expressed from a human cytomegalovirus immediate early promoter. Recombinant HVT expressing cIFN is useful as a vaccine in poultry against Marek's Disease.

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Recombinant HVT expressing cIFN is constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes were used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 761-07.A1 with Asc I, 415-09.BA1 with BamHI.

Recombinant HVT expressing avian cytokines is combined with HVT expressing genes for avian disease antigens to enhance immune response. Additional cytokines that are expressed in HVT and have immune stimulating effects include, but not limited to, transforming growth factor beta, epidermal growth factor family, fibroblast growth factors, hepatocyte growth factor, insulin-like growth factors, B-nerve growth factor, platelet-derived growth factor, vascular endothelial growth factor, interleukin 1, IL-1 receptor antagonist, interleukin 2, interleukin 3, interleukin 4, interleukin 5, interleukin 6, IL-6 interleukin 7, soluble receptor, interleukin 8, interleukin 9, interleukin 10, interleukin interleukin 12, interleukin 13, angiogenin, chemokines, colony stimulating factors, granulocyte-macrophage colony stimulating factors, erythropoietin, interferon, interferon gamma, leukemia inhibitory factor, oncostatin Μ, pleiotrophin, secretory leukocyte protease inhibitor, stem cell factor, tumor necrosis factors, and soluble TNF receptors. These cytokines are

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from avian species or other animals including humans, bovine, equine, feline, canine or porcine.

Example 20C Recombinant HVT expressing Marek's disease virus genes and chicken interferon gene.

A recombinant herpesvirus of turkeys contains the chicken interferon (cIFN) gene inserted into an Xhol site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT and further contains the MDV gA, gD, and gB genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The cIFN gene is expressed from an human cytomegalovirus immediate early promoter. The MDV genes are expressed from the endogenous MDV promoters. Recombinant HVT expressing cIFN and MDV gA, gB, and gD is useful as a vaccine with an enhanced immune response in poultry against Marek's Disease.

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Recombinant HVT expressing MDV genes and the cIFN gene is constructed according to the PROCEDURE FROM GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes are used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 761-07.A1 with Asc I, 721-38.1J uncut.

30 <u>Example 20D</u> Recombinant HVT expressing Marek's disease virus genes, Newcastle disease virus genes and chicken interferon gene.

A recombinant herpesvirus of turkeys contains the chicken interferon (cIFN) gene inserted into an XhoI site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT and further

contains the MDV gA, gD, and gB genes and NDV HN and F genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The cIFN gene is expressed from an human cytomegalovirus immediate early promoter. The MDV genes are expressed from the endogenous MDV promoters. The NDV HN gene is under the control of the PRV gX promoter, and the NDV F gene is under the control of the HCMV immediate early promoter. Recombinant HVT expressing cIFN and MDV gA, gB, and gD is useful as a vaccine with an enhanced immune response in poultry against Marek's Disease and Newcastle disease.

Recombinant HVT expression MDV genes, NDV genes and cIFN is constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes are used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 761-07.A1 with Asc I, 722-60.E2 uncut.

Example 20E Recombinant HVT expressing Marek's disease virus genes and chicken myelomonocytic growth factor gene.

A recombinant herpesvirus of turkeys contains the chicken myelomonocytic growth factor (cMGF) gene inserted into and XhoI site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT and further contains the MDV gA, gD, and gB genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. The cMGF gene is expressed from a human cytomegalovirus immediate early promoter. The MDV genes are expressed from the endogenous MDV promoters. Recombinant HVT expression cMGF and MDV gA, gB, and gD is useful as a vaccine with

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an enhanced immune response in poultry\_against Marek's Disease.

Recombinant HVT expressing the cMGF gene and MDV genes is constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes are used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 751-87.A8 with Asc I, 721-38.1J uncut.

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Example 20F Recombinant HVT expressing Marek's disease virus genes, Newcastle disease virus genes and chicken myelomonocytic growth factor gene.

A recombinant herpesvirus of turkeys contains the chicken myelomonocytic growth factor (cGMF) inserted into an XhoI site converted to a PacI site in the EcoR1 #9 fragment within the unique long region of HVT and further contains the MDV gA, gD, and gB genes and NDV HN and F genes inserted into a unique StuI site converted into a HindIII site in the HVT US2 gene. cGMF gene is expressed from an human cytomegalovirus immediate early promoter. The MDV genes are expressed from the endogenous MDV promoters. The NDV HN gene is under the control of the PRV gX promoter, and the NDV F gene is under the control of the HCMV immediate early promoter. Recombinant HVT expressing cIFN and MDV gA, gB and gD is useful as a vaccine with an enhanced immune response in poultry against Marek's Disease and Newcastle disease.

Recombinant HVT expressing MDV genes, NDV genes and the cGMF gene is constructed according to the PROCEDURE FOR GENERATING RECOMBINANT HERPESVIRUS FROM OVERLAPPING

SUBGENOMIC FRAGMENTS. The following combination of subgenomic clones and enzymes are used: 407-32.2C3 with NotI, 172-07.BA2 with BamHI, 407-32.5G6 with NotI, 672-07.C40 with NotI, 672-01.A40 with NotI, 751-87.A8 uncut, 722-60.E2 uncut.

Example 21 Recombinant herpesvirus of turkeys expressing antigens from disease causing microorganisms

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Recombinant herpesvirus of turkeys (HVT) is useful for expressing antigens from disease causing microorganisms from animals in addition to avian species. Recombinant HVT is useful as a vaccine in animals including but not limited to humans, equine, bovine, porcine, canine and feline.

Recombinant HVT is useful as a vaccine against equine diseases when foreign antigens from diseases or disease organisms are expressed in the HVT vector, including not limited to: eguine influenza, herpesvirus-1 and equine herpesvirus-4. Recombinant HVT is useful as a vaccine against bovine diseases when foreign antigens from the following diseases or disease organisms are expressed in the HVT vector, including, but not limited to: bovine herpesvirus type 1, bovine viral diarrhea virus, bovine respiratory syncytial virus, bovine parainfluenza virus. Recombinant HVT is useful as a vaccine against swine diseases when foreign antigens from the following diseases or disease organisms are expressed in the HVT vector, including limited to: pseudorabies virus, but not reproductive and respiratory syndrome (PRRS/SIRS), hog cholera virus, swine influenza virus, swine parvovirus, swine rotavirus. Recombinant HVT is useful as a vaccine against feline or canine diseases when foreign antigens from the following diseases or disease organisms are

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expressed in the HVT vector, including but not limited to feline herpesvirus, feline leukemia virus, feline immunodeficiency virus and Dirofilaria (heartworm). Disease causing microorganisms in dogs include, but are not limited to canine herpesvirus, canine distemper, canine adenovirus type 1 (hepatitis), adenovirus type 2 (respiratory disease), parainfluenza, Leptospira canicola, icterohemorragia, parvovirus, coronavirus, Borrelia burgdorferi, canine herpesvirus, Bordetella bronchiseptica, Dirofilaria immitis (heartworm) and rabies virus.

Example 22 Human vaccines using recombinant herpesvirus of turkeys as a vector

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Recombinant herpesvirus of turkeys (HVT) is useful as a vaccine against human diseases. For example, human influenza is a rapidly evolving virus neutralizing viral epitopes are rapidly changing. A useful recombinant HVT vaccine is one in which the influenza neutralizing epitopes are quickly changed to protect against new strains of influenza. influenza HA and NA genes are cloned using polymerase chain reaction into the recombinant HVT. Recombinant HVT is useful as a vaccine against other human diseases when foreign antigens from the following diseases or disease organisms are expressed in the HVT vector: hepatitis B virus surface and core antigens, hepatitis C virus, human immunodeficiency virus, herpes simplex virus-1, herpes simplex virus-2, human cytomegalovirus, Epstein-Barr virus, Varicella-Zoster virus, herpesvirus-6, human herpesvirus-7, human influenza, measles virus, hantaan virus, pneumonia rhinovirus, poliovirus, human respiratory syncytial virus, retrovirus, human T-cell leukemia virus, rabies virus, mumps virus, malaria (Plasmodium falciparum), Bordetella pertussis, Diptheria, Rickettsia prowazekii,

Borrelia bergdorferi, Tetanus toxoid, malignant tumor antigens,

Recombinant HVT expressing human cytokines is combined with HVT expressing genes for human disease antigens to enhance immune response. Additional cytokines, including, limited to, transforming but not factor beta, epidermal growth factor family, fibroblast growth factors, hepatocyte growth factor, insulin-like growth factors, B-nerve growth factor, platelet-derived growth factor, vascular endothelial growth factor, interleukin 1, IL-1 receptor antagonist, interleukin 2, interleukin 3, interleukin 4, interleukin 5, interleukin 6, IL-6 soluble receptor, interleukin 7, interleukin 8, interleukin 9, interleukin interleukin 11, interleukin 12, interleukin 13, angiogenin, chemokines, colony stimulating factors, granulocyte-macrophage colony stimulating factors, erythropoietin, interferon, interferon gamma, leukemia inhibitory factor, oncostatin Μ, pleiotrophin, secretory leukocyte protease inhibitor, stem cell tumor necrosis factors, and soluble receptors from human and other animals are expressed in HVT and have immune stimulating effects.

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Example 23 Improved production of a recombinant herpesvirus of turkeys vaccine.

Cytokines, such as interferons and interleukins, inhibit the replication of viruses in cell culture and in the animal. Inhibition of the production of cellular interferon or interleukin improves the growth of recombinant HVT in cell culture. Chicken interferon (cIFN) expressed from a recombinant swinepox vector was added to chick embryo fibroblast (CEF) cell cultures and infected with S-HVT-012 which expresses ß-galactosidase. cIFN added to the cell culture media

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reduced both the expression of ß-galactosidase and S-HVT-012 titer in a dose dependent manner. This result indicates that growth of HVT is limited by exogenous addition of chicken interferon. Several strategies are utilized to improve growth of HVT in CEF cells by removing or inactivating chicken interferon activity in the CEF cells.

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In one embodiment, a chicken interferon neutralizing antibody is added to the culture medium to inhibit the chicken interferon activity and improve the growth of recombinant HVT in CEF cell culture. The anti-cIFN antibody is derived from mouse or rabbit sera of animals injected with chicken interferon protein, preferably the cIFN is from a recombinant swinepox virus expressing chicken interferon.

Poxviruses secrete cytokine-inhibiting proteins as an immune evasion strategy. One type of poxvirus immune evasion mechanism involves poxvirus soluble receptors for interleukins, interferon, or tumor necrosis factors cytokines and allow inactive the replication (60). In an embodiment of the invention, fowlpox virus is useful as a source of chicken interferon-inhibiting proteins and other immune evasion proteins. Conditioned media from FPV infected CEF cell cultures is added to the HVT infected CEF cells to inhibit interferon activity and increase the HVT titer. In a further embodiment, the recombinant chicken interferon inhibiting protein or another poxvirus immune evasion protein is expressed in a vector in combination with an HVT vaccine composition to increase the HVT titer.

Chicken embryo fibroblast cells have been engineered to express foreign genes (61). in a further embodiment, an interferon-negative CEF cell line is constructed by

the introduction of a vector expressing a gene encoding antisense RNA for chicken interferon into the CEF cell line. Recombinant HVT grown in an interferon-negative CEF cell line demonstrate improved virus titers compared to HVT grown in an interferon producing CEF In a further embodiment, a chicken myelomonocytic growth factor (cMGF) -positive CEF cell line is constructed by the introduction of a vector CEF cells. expressing the cMGF gene into the Recombinant HVT grown in a cMGF-positive CEF cell line demonstrates improved virus titers compared to HVT grown in a cMGF negative CEF cell line.

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Recombinant HVT of the present invention is useful as
a vaccine against Marek's disease and against other
diseases as outlined in previous examples. An
increased efficiency in growth of recombinant HVT in
CEF cells is useful in production of the vaccine.

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<ul> <li>(iv) CORRESPONDENCE ADDRESS:</li> <li>(A) ADDRESSEE: John P. White</li> <li>(B) STREET: 1185 Avenue of the Americas</li> <li>(C) CITY: New York</li> <li>(D) STATE: New York</li> <li>(E) COUNTRY: USA</li> <li>(F) ZIP: 10036</li> </ul>
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(2) INFORMATION FOR SEQ ID NO:1:
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 3350 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: DNA (genomic)
(iii) HYPOTHETICAL: NO
(iv) ANTI-SENSE: NO
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1292522
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
GGATACGATC GGTCTGACCC GGGGGAGTCA CCCGGGGACA GCCGTCAAGG CCTTGTTCCA 60
GGATAGAACT CCTCCTTCTA CAACGCTATC ATTGATGGTC AGTAGAGATC AGACAAACGA 120

TCGCAGCG ATG ACA AAC CTG CAA GAT CAA ACC CAA CAG ATT GTT CCG TTC Met Thr Asn Leu Gln Asp Gln Thr Gln Gln Ile Val Pro Phe 1 5 10

ATA Ile 15	CGG Arg	AGC Ser	CTT Leu	CTG Leu	ATG Met 20	CCA Pro	ACA Thr	ACC Thr	GGA Gly	CCG Pro 25	GCG Ala	TCC Ser	ATT Ile	CCG Pro	GAG Glu 30	218	
ACA Thr	CCC Pro	TGG Trp	AGA Arg	AGC Ser 35	ACA Thr	CTC Leu	TCA Ser	GGT Gly	CAG Gln 40	AGA Arg	CTG Leu	ACC Thr	TAC Tyr	AAT Asn 45	TTG Leu	266	
ACT Thr	GTG Val	GGG Gly	GAC Asp 50	ACA Thr	GGG Gly	TCA Ser	GGG Gly	CTA Leu 55	ATT Ile	GTC Val	TTT Phe	TTC Phe	CCT Pro 60	GGA Gly	TTC Phe	314	
CCT Pro	GGC Gly	TCA Ser 65	ATT Ile	GTG Val	GGT Gly	GCT Ala	CAC His 70	TAC Tyr	ACA Thr	CTG Leu	CAG Gln	AGC Ser 75	AAT Asn	GGG Gly	AAC Asn	362	
TAC Tyr	AAG Lys 80	TTC Phe	GAT Asp	CGG Arg	ATG Met	CTC Leu 85	CTG Leu	ACT Thr	GCC Ala	CAG Gln	AAC Asn 90	CTA Leu	CCG Pro	GCC Ala	AGT Ser	410	
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ACA Thr	CTT Leu	CCT Pro	GGT Gly	GGC Gly 115	GTT Val	TAT Tyr	GCA Ala	CTA Leu	AAC Asn 120	GGC Gly	ACC Thr	ATA Ile	AAC Asn	GCC Ala 125	GTG Val	506	
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GTA Val	GGG Gly 160	GAA Glu	GGG Gly	GTC Val	ACC Thr	GTC Val 165	CTC Leu	AGC Ser	TTA Leu	CCC Pro	ACA Thr 170	TCA Ser	TAT Tyr	GAT Asp	CTT Leu	650	
GGG Gly 175	TAT Tyr	GTG Val	AGG Arg	CTT Leu	GGT Gly 180	GAC Asp	CCC Pro	ATT Ile	CCC Pro	GCA Ala 185	ATA Ile	GGG Gly	CTT Leu	GAC Asp	CCA Pro 190	698	
AAA Lys	ATG Met	GTA Val	GCC Ala	ACA Thr 195	TGT Cys	GAC Asp	AGC Ser	AGT Ser	GAC Asp 200	AGG Arg	CCC Pro	AGA Arg	GTC Val	TAC Tyr 205	ACC Thr	746	
ATA Ile	ACT Thr	GCA Ala	GCC Ala 210	Asp	GAT Asp	TAC Tyr	CAA Gln	TTC Phe 215	ser	TCA Ser	CAG Gln	TAC Tyr	CAA Gln 220	CCA Pro	GGT Gly	794	
GGG Gly	GTA Val	ACA Thr 225	Ile	ACA Thr	CTG Leu	TTC Phe	TCA Ser 230	Ala	AAC Asn	ATT Ile	GAT Asp	GCC Ala 235	ATC Ile	ACA Thr	AGC Ser	842	
CTC Leu	AGC Ser 240	Val	GGG Gly	GGA Gly	GAG Glu	CTC Leu 245	Val	TTT Phe	CGA Arg	ACA Thr	AGC Ser 250	Val	CAC	GGC	CTT Leu	890	I
GTA Val 255	Leu	GGC Gly	GCC Ala	ACC Thr	Ile 260	Tyr	CTC Leu	ATA	GGC Gly	TTT Phe 265	Asp	GGG Gly	ACA Thr	ACG Thr	GTA Val 270	938	ļ
ATC Ile	ACC Thr	AGG Arg	GCT Ala	GTG Val	Ala	GCA Ala	AAC Asn	ACT Thr	GGG Gly 280	Leu	ACG Thr	ACC Thr	GGC Gly	ACC Thr 285	GAC Asp	986	;

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CCA Pro	ATC Ile	ACA Thr 305	TCC Ser	ATC Ile	AAA Lys	CTG Leu	GAG Glu 310	ATA Ile	GTG Val	ACC Thr	TCC Ser	AAA Lys 315	AGT Ser	GGT Gly	GGT	1082
										AGA Arg		Ser				1130
										CTC Leu 345						1178
										GTC Val						1226
										GAA Glu						1274
Val	Thr	Glu 385	Tyr	Gly	Arg	Phe	Asp 390	Pro	Gly	GCC Ala	Met	Asn 395	Tyr	Thr	Lys	1322
Leu	Ile 400	Leu	Ser	Glu	Arg	Asp 405	Arg	Leu	Gly	ATC Ile	Lys 410	Thr	Val	Trp	Pro	1370
Thr 415	Arg	Glu	Tyr	Thr	Asp 420	Phe	Arg	Glu	Tyr	TTC Phe 425	Met	Glu	Val	Ala	Asp 430	1418
Leu	Asn	Ser	Pro	Leu 435	Lys	Ile	Ala	Gly	Ala 440	TTC Phe	Gly	Phe	Lys	<b>Asp</b> 445	Ile	1466
Ile	Arg	Ala	Ile 450	Arg	Arg	Ile	Ala	Val 455	Pro	GTG Val	Val	Ser	Thr 460	Leu	Phe	1514
Pro	Pro	Ala 465	Ala	Pro	Leu	Ala	His 470	Ala	Ile	GGG Gly	Glu	Gly 475	Val	Asp	Tyr	1562
Leu	Leu 480	Gly	Asp	Glu	Ala	Gln 485	Ala	Ala	Ser	GGA Gly	Thr 490	Ala	Arg	Ala	Ala	1610
Ser 495	Gly	Lys	Ala	Arg	Ala 500	Ala	Ser	Gly	Arg	ATA Ile 505	Arg	Gln	Leu	Thr	Le <b>u</b> 510	1658
Ala	Ala	Asp	Lys	Gly 515	Tyr	Glu	Val	Val	Ala 520	AAT Asn	Leu	Phe	Gln	Val 525	Pro	1706
Gln	Asn	Pro	Val 530	Val	Asp	Gly	Ile	Leu 535	Ala	TCA Ser	Pro	Gly	Val 540	Leu	Arg	1754
GGT Gly	GCA Ala	CAC His 545	AAC Asn	CTC Leu	GAC Asp	TGC Cys	GTG Val 550	TTA Leu	AGA Arg	GAG Glu	GGT Gly	GCC Ala 555	ACG Thr	CTA Leu	TTC Phe	1802

CCT Pro	GTG Val 560	Val	ATT Ile	ACG Thr	ACA Thr	G <b>TG</b> Val 565	GAA Glu	GAC Asp	GCC Ala	ATG Met	ACA Thr 570	CCC Pro	AAA Lys	GCA Ala	TTG Leu	1850
AAC Asn 575	Ser	AAA Lys	ATG Met	TTT Phe	GCT Ala 580	GTC Val	ATT Ile	GAA Glu	GGC Gly	GTG Val 585	CGA Arg	GAA Glu	GAC Asp	CTC Leu	CAA Gln 590	1898
CCT Pro	CCA Pro	TCT Ser	CAA Gln	AGA Arg 595	G <b>GA</b> Gly	TCC Ser	TTC Phe	ATA Ile	CGA Arg 600	ACT Thr	CTC Leu	TCT Ser	GGA Gly	CAC His 605	AGA Arg	1946
GTC Val	TAT Tyr	GGA Gly	TAT Tyr 610	GCT Ala	CCA Pro	GAT Asp	GG Gly	GTA Val 615	CT <b>T</b> Leu	CCA Pro	CTG Leu	GAG Glu	ACT Thr 620	GGG Gly	AGA Arg	1994
GAC Asp	TAC Tyr	ACC Thr 625	GTT Val	GTC Val	CCA Pro	ATA Ile	GAT Asp 630	GAT Asp	GTC Val	TGG Trp	GAC Asp	GAC Asp 635	AGC Ser	ATT Ile	ATG Met	2042
CTG Leu	TCC Ser 640	AAA Lys	GAT Asp	CCC Pro	ATA Ile	CCT Pro 645	CCT Pro	ATT Ile	GTG Val	GGA Gly	AAC Asn 650	AGT Ser	GGA Gly	AAT Asn	CTA Leu	2090
GCC Ala 655	Ile	GCT Ala	TAC Tyr	ATG Met	GAT Asp 660	GTG Val	TTT Phe	CGA Arg	CCC Pro	AAA Lys 665	Val	CCA Pro	ATC Ile	CAT His	GTG Val 670	2138
GCT Ala	ATG Met	ACG Thr	GGA Gly	GCC Ala 675	CTC Leu	AAT Asn	GCT Ala	TGT Cys	GGC Gly 680	GAG Glu	ATT Ile	GAG Glu	AAA Lys	GTA Val 685	AGC Ser	2186
TTT Phe	AGA Arg	AGC Ser	ACC Thr 690	AAG Lys	CTC Leu	GCC Ala	ACT Thr	GCA Ala 695	CAC His	CGA Arg	C <b>T</b> T Leu	GGC Gly	CTT Leu 700	AAG Lys	TTG Leu	2234
GCT Ala	GGT Gly	CCC Pro 705	GGA Gly	GCA Ala	TTC Phe	GAT Asp	GTA Val 710	AAC Asn	ACC Thr	GGG Gly	CCC Pro	AAC Asn 715	TGG Trp	GCA Ala	ACG Thr	2282
TTC Phe	ATC Ile 720	AAA Lys	CGT <b>A</b> rg	TTC Phe	CCT Pro	CAC His 725	AAT <b>A</b> sn	CCA Pro	CGC Arg	GAC Asp	TGG Trp 730	GAC Asp	AGG Arg	CTC <b>Leu</b>	CCC Pro	2330
						CTT Leu										2378
CTT Leu	GCC Ala	ATG Met	GCT Ala	GCA Ala 755	TCA Ser	GAG Glu	TTC Phe	AAG Lys	AGA Arg 760	CCC Pro	CGA Arg	ACT Thr	CGA Arg	GAG Glu 765	TGC Cys	2426
CGT <b>Ar</b> g						AGC Ser										2474
TGC Cys						Val									TGA	2522
CATG	GCCA	AC T	TCGC	ACTC	A GC	GACC	CGAA	CGC	CCAT	CGG	ATGC	GAAA	TT I	TTTI	'GCAAA	2582
CGAC	CACA	AG C	AGGC	AGCA	A GT	CGCA	AAGG	GCC	AAGT	ACG	GGAC	AGCA	.GG C	TACG	GAGTG	2642
GAGG	CTCG	GG G	cccc	CACA	C CA	GAGG.	AAGC	ACA	GAGG	GAA	AAAG	ACAC	AC G	GATC	TCAAA	2702
GAAG	ATGG	AG A	CCAT	GGGC	A TC	TACT	TTGC	AAC	ACCA	GAA	TGGG	TAGC	AC T	CAAT	GGGCA	2762

CCGAGGGCCA	AGCCCCGGCC	AGCTAAAGTA	CGGGCAGAAC	ACACGAGAAA	TACGGACCCA	2822
AACGAGGACT	ATCTAGACTA	CGTGCATGCA	GAGAAGAGCC	GGTTGGCATC	AGAAGAACAA	2882
ATCCTAAGGG	CAGCTACGTC	AGATCTACGG	GGCTCCAGGA	CAGGCAGAGC	ACCCCAAGCT	2942
TTCATAGACG	AAGTTGCCAA	AGTCTATGAA	ATCAACCATG	GACGTGGCCC	AAACCAAGAA	3002
CAGATGAAAG	ATCTGCTCTT	GACTGCGATG	GAGATGAAGC	ATCGCAATCC	CAGGCGGGCT	3062
CTACCAAAGC	CCAAGCCAAA	ACCCAATGCT	CCAACACAGA	GACCCCCTGG	TCGGCTGGGG	3122
CTGGATCAGG	ACCGTCTCTG	ATGAGGACCT	TGAGTGAGGC	TCCTGGGAGT	CTCCCGACAA	3182
CACCCGCGCA	GGTGTGGACA	CAATTCGGCC	TTACAACATC	CCAAATTGGA	TCCGTTCGCG	3242
GGTCCCCAAA	ААААААААА	АААААААА	AAAAAAAA	АААААААА	AAAAAAAA	3302
AAGTACCTTC	TGAGGCGGAA	AGAACCAGCC	GGATCCCTCG	AGGGATCC		3350

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 797 amino acids
  - (B) TYPE: amino acid(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Thr Asn Leu Gln Asp Gln Thr Gln Gln Ile Val Pro Phe Ile Arg

Ser Leu Leu Met Pro Thr Thr Gly Pro Ala Ser Ile Pro Glu Thr Pro

Trp Arg Ser Thr Leu Ser Gly Gln Arg Leu Thr Tyr Asn Leu Thr Val

Gly Asp Thr Gly Ser Gly Leu Ile Val Phe Phe Pro Gly Phe Pro Gly

Ser Ile Val Gly Ala His Tyr Thr Leu Gln Ser Asn Gly Asn Tyr Lys

Phe Asp Arg Met Leu Leu Thr Ala Gln Asn Leu Pro Ala Ser Tyr Asn 90

Tyr Cys Arg Leu Val Ser Arg Ser Leu Thr Val Arg Ser Ser Thr Leu

Pro Gly Gly Val Tyr Ala Leu Asn Gly Thr Ile Asn Ala Val Thr Phe 115

Gln Gly Ser Leu Ser Glu Leu Thr Asp Val Ser Tyr Asn Gly Leu Met 135

Ser Ala Thr Ala Asn Ile Asn Asp Lys Ile Gly Asn Val Leu Val Gly 145 155

Glu Gly Val Thr Val Leu Ser Leu Pro Thr Ser Tyr Asp Leu Gly Tyr

Val Arg Leu Gly Asp Pro Ile Pro Ala Ile Gly Leu Asp Pro Lys Met

			180					185					190		
Val	Ala	Thr 195	Cys	Asp	Ser	Ser	Asp 200	Arg	Pro	Arg	Val	Tyr 205	Thr	Ile	Thr
Ala	Ala 210	Asp	Asp	Tyr	Gln	Phe 215	Ser	Ser	Gln	Tyr	Gln 220	Pro	Gly	Gly	Val
Thr 225	Ile	Thr	Leu	Phe	Ser 230	Ala	Asn	Ile	Asp	Ala 235	Ile	Thr	Ser	Leu	Ser 240
Val	Gly	Gly	Glu	Leu 245	Val	Phe	Arg	Thr	Ser 250	Val	His	Gly	Leu	Val 255	Leu
Gly	Ala	Thr	Ile 260	Tyr	Leu	Ile	Gly	Phe 265	Asp	Gly	Thr	Thr	Val 270	Ile	Thr
Arg	Ala	Val 275	Ala	Ala	Asn	Thr	Gly 280	Leu	Thr	Thr	Gly	Thr 285	Asp	Asn	Leu
Met	Pro 290	Phe	Asn	Leu	Val	Ile 295	Pro	Thr	Asn	Glu	Ile 300	Thr	Gln	Pro	Ile
Thr 305	Ser	Ile	Lys	Leu	Glu 310	Ile	Val	Thr	Ser	Lys 315	Ser	Gly	Gly	Gln	Ala 320
Gly	Asp	Gln	Met	Leu 325	Trp	Ser	Ala	Arg	Gly 330	Ser	Leu	Ala	Val	Thr 335	Ile
His	Gly	Gly	Asn 340	Tyr	Pro	Gly	Ala	Leu 345	Arg	Pro	Val	Thr	Leu 350	Val	Ala
Tyr	Glu	Arg 355	Val	Ala	Thr	Gly	Ser 360	Val	Val	Thr	Val	Ala 365	Gly	Val	Ser
Asn	Phe 370	Glu	Leu	Ile	Pro	Asn 375	Pro	Glu	Leu	Ala	Lys 380	Asn	Leu	Val	Thr
Glu 385	Tyr	Gly	Arg	Phe	Asp 390	Pro	Gly	Ala	Met	Asn 395	Tyr	Thr	Lys	Leu	Ile 400
Leu	Ser	Glu	Arg	Asp 405	Arg	Leu	Gly	Ile	Lys 410	Thr	Val	Trp	Pro	Thr 415	Arg
Glu	Tyr	Thr	Asp 420	Phe	Arg	Glu	Tyr	Phe 425	Met	Glu	Val	Ala	Asp 430	Leu	Asn
Ser	Pro	Leu 435	Lys	Ile	Ala	Gly	Ala 440	Phe	Gly	Phe	Lys	Asp 445	Ile	Ile	Arg
Ala	Ile 450	Arg	Arg	Ile	Ala	Val 455	Pro	Val	Val	Ser	Thr 460	Leu	Phe	Pro	Pro
Ala 465	Ala	Pro	Leu	Ala	His 470	Ala	Ile	Gly	Glu	Gly 475	Val	Asp	Tyr	Leu	Leu 480
Gly	Asp	Glu	Ala	Gln 485	Ala	Ala	Ser	Gly	Thr 490	Ala	Arg	Ala	Ala	Ser 495	Gly
Lys	Ala	Arg	Ala 500	Ala	Ser	Gly	Arg	Ile 505	Arg	Gln	Leu	Thr	Leu 510	Ala	Ala
Asp	Lys	Gly 515	Tyr	Glu	Val	Val	Ala 520	Asn	Leu	Phe	Gln	Val 525	Pro	Gln	Asn
Pro	Val	Val	Asp	Gly	Ile	Leu	Ala	Ser	Pro	Gly	Val	Leu	Arg	Gly	Ala

530 535 540

His Asn Leu Asp Cys Val Leu Arg Glu Gly Ala Thr Leu Phe Pro Val 545 550 555 560

Val Ile Thr Thr Val Glu Asp Ala Met Thr Pro Lys Ala Leu Asn Ser 565 570 575

Lys Met Phe Ala Val Ile Glu Gly Val Arg Glu Asp Leu Gln Pro Pro 580 585 590

Ser Gln Arg Gly Ser Phe Ile Arg Thr Leu Ser Gly His Arg Val Tyr 595 600 605

Gly Tyr Ala Pro Asp Gly Val Leu Pro Leu Glu Thr Gly Arg Asp Tyr 610 625

Thr Val Val Pro Ile Asp Asp Val Trp Asp Asp Ser Ile Met Leu Ser 625 630 635 640

Lys Asp Pro Ile Pro Pro Ile Val Gly Asn Ser Gly Asn Leu Ala Ile 645 650 655

Ala Tyr Met Asp Val Phe Arg Pro Lys Val Pro Ile His Val Ala Met 660 665 670

Thr Gly Ala Leu Asn Ala Cys Gly Glu Ile Glu Lys Val Ser Phe Arg 675 680 685

Ser Thr Lys Leu Ala Thr Ala His Arg Leu Gly Leu Lys Leu Ala Gly 690 695 700

Pro Gly Ala Phe Asp Val Asn Thr Gly Pro Asn Trp Ala Thr Phe Ile 705 710 715 720

Lys Arg Phe Pro His Asn Pro Arg Asp Trp Asp Arg Leu Pro Tyr Leu 725 730 735

Asn Leu Pro Tyr Leu Pro Pro Asn Ala Gly Arg Gln Tyr His Leu Ala
740 745 750

Met Ala Ala Ser Glu Phe Lys Arg Pro Arg Thr Arg Glu Cys Arg Gln 755 760 765

Ser Asn Gly Ser Ser Ser Gln Arg Gly Pro Thr Ile Pro Ile Cys Thr 770 775 780

Gln Cys Val His Val Ala Gly Arg Glu Trp Asp Cys Asp
785 790 795

### (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 5426 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (ix) FEATURE:
  - (A) NAME/KEY: CDS

157

(B) LOCATION: 73..1182

		(	D) O	THER	INF	ORMA	TION	: /p	rodu	ct=	"HVT	UL4	2"			
	(ix	(	B) L	AME/ OCAT	KEY: ION: INF	130	62		rodu	ct=	"HVT	UL4	3"			
	(ix	{	B) L	AME/	KEY: ION: INF	279	04		rodu	ct=	"HVT	gA"				
	(ix	(. (:	B) L	AME/	KEY: ION: INF	470			rodu	ct=	"HVT	UL4	5 <b>"</b>			
	(xi	) SE	QUEN	CE D	ESCR:	IPTI	ON:	SEQ :	ID N	0:3:						
GGAT	rccg/	AGC '	TTCT	ACTA'	TA C	AACG	CGGA	C GA	TAAT'	TTTG	TCC	ACCC	CAT	CGGT	G <b>T</b> TCGA	60
GAA	AGGG?	<b>ГТ</b> Т '			TG G( et A											108
GCA Ala	GGA Gly	GAG Glu 15	GCT Ala	CAT His	ACA Thr	CCC Pro	GAG Glu 20	GAT Asp	ATG Met	CAA Gln	AAG Lys	AAA Lys 25	TGG Trp	AGG Arg	ATT Ile	156
ATA Ile	TTG Leu 30	GCA Ala	GGG Gly	GAA Glu	AAA Lys	TTC Phe 35	ATG Met	ACT Thr	ATA Ile	TCG Ser	GCA Ala 40	TCG Ser	TTG Leu	AAA Lys	TCG Ser	204
					AAA Lys 50											252
					ACT Thr											300
					AGC Ser											348
					GAT Asp											396
					GCA Ala											444
CCT Pro 125	CCA Pro	TGT Cys	AGG Arg	CAT His	TTA Leu 130	ATC Ile	CAA Gln	GCC Ala	GTC Val	ACA Thr 135	TAC Tyr	ATG Met	ACC Thr	GAC Asp	GGT Gly 140	492
					ACA Thr											5 <b>4</b> 0
					AAA Lys											588

CAA CAA TTG AAC AAA ATA TTG GCC GTC GCT TCA AAA CTG CAA CAC GAA Gln Gln Leu Asn Lys Ile Leu Ala Val Ala Ser Lys Leu Gln His Glu 175 180 185	636
GAA CTT GTA TTC TCT TTA AAA CCT GAA GGA GGG TTC TAC GTA GGA ACG Glu Leu Val Phe Ser Leu Lys Pro Glu Gly Gly Phe Tyr Val Gly Thr 190 195 200	684
GTT TGT ACT GTT ATA AGT TTC GAA GTA GAT GGG ACT GCC ATG ACT CAG Val Cys Thr Val Ile Ser Phe Glu Val Asp Gly Thr Ala Met Thr Gln 205 210 215 220	732
TAT CCT TAC AAC CCT CCA ACC TCG GCT ACC CTA GCT CTC GTA GCA Tyr Pro Tyr Asn Pro Pro Thr Ser Ala Thr Leu Ala Leu Val Val Ala 225 230 235	780
TGC AGA AAG AAG AAG GCG AAT AAA AAC ACT ATT TTA ACG GCC TAT GGA Cys Arg Lys Lys Ala Asn Lys Asn Thr Ile Leu Thr Ala Tyr Gly 240 245 250	828
AGT GGT AAA CCC TTT TGT GTT GCA TTG GAA GAT ACT AGT GCA TTT AGA Ser Gly Lys Pro Phe Cys Val Ala Leu Glu Asp Thr Ser Ala Phe Arg 255 260 265	876
AAT ATC GTC AAT AAA ATC AAG GCG GGT ACG TCG GGA GTT GAT CTG GGG Asn Ile Val Asn Lys Ile Lys Ala Gly Thr Ser Gly Val Asp Leu Gly 270 . 275 280	924
TTT TAT ACA ACT TGC GAT CCG CCG ATG CTA TGT ATT CGC CCA CAC GCA Phe Tyr Thr Thr Cys Asp Pro Pro Met Leu Cys Ile Arg Pro His Ala 285 290 295 300	972
TTT GGA AGT CCT ACC GCA TTC CTG TTT TGT AAC ACA GAC TGT ATG ACA Phe Gly Ser Pro Thr Ala Phe Leu Phe Cys Asn Thr Asp Cys Met Thr 305 310 315	1020
ATA TAT GAA CTG GAA GAA GTA AGC GCC GTT GAT GGT GCA ATC CGA GCA Ile Tyr Glu Leu Glu Glu Val Ser Ala Val Asp Gly Ala Ile Arg Ala 320 -325 330	1068
AAA CGC ATC AAC GAA TAT TTC CCA ACA GTA TCG CAG GCT ACT TCC AAG Lys Arg Ile Asn Glu Tyr Phe Pro Thr Val Ser Gln Ala Thr Ser Lys 335 340 345	1116
AAG AGA AAA CAG TCG CCG CCC CCT ATC GAA AGA GAA AGG AAA ACC ACC Lys Arg Lys Gln Ser Pro Pro Pro Ile Glu Arg Glu Arg Lys Thr Thr 350 355 360	1164
AGA GCG GAT ACC CAA TAAAATGCCA GACAAACCCG GCATCCTGGT TAGAGGGCAG Arg Ala Asp Thr Gln 365 370	1219
GTGGGCTGGG CCAACCTTCA CGGGCGTCCG ACAGATCGGT GACACTCATA CGTTAACTAA	1279
ACGCCGGCAG CTTTGCAGAA GAAAAT ATG CCT TCC GGA GCC AGC TCG AGT CCT  Met Pro Ser Gly Ala Ser Ser Pro  1 5	1332
CCA CCA GCT TAT ACA TCT GCA GCT CCG CTT GAG ACT TAT AAC AGC TGG Pro Pro Ala Tyr Thr Ser Ala Ala Pro Leu Glu Thr Tyr Asn Ser Trp 10 15 20 25	1380
CTA AGT GCC TTT TCA TGC GCA TAT CCC CAA TGC ACT GCG GGA AGA GGA Leu Ser Ala Phe Ser Cys Ala Tyr Pro Gln Cys Thr Ala Gly Arg Gly 30 35 40	1428

	CGA Arg															1476
TGT Cys	TCC Ser	TTA Leu 60	GTG Val	TGC Cys	ATC Ile	GCT Ala	GCA Ala 65	CAT His	TTA Leu	GCT Ala	GTT Val	ACC Thr 70	GTG Val	TCG Ser	GGA Gly	1524
	GCA Ala 75															1572
	GTA Val															1620
	ATA Ile															1668
	ATA Ile					-	_								_	1716
	GCG Ala															1764
	GAA Glu 155		_													1812
	TCC Ser															1860
AAG Lys	GGC Gly	GGG Gly	ATT Ile	TTT Phe 190	CAT His	GCT Ala	TAC Tyr	CAC His	GGA Gly 195	ACA Thr	TTA Leu	CTC Leu	GGT Gly	ATA Ile 200	ACA Thr	1908
ATA Ile	CCA Pro	AAC Asn	ATA Ile 205	CAC His	CCA Pro	ATC Ile	CCT Pro	CTC Leu 210	GCG Ala	GGG Gly	TTT Phe	CTT Leu	GCA Ala 215	GTC Val	TAT Tyr	1956
ACA Thr	ATA Ile	TTG Leu 220	GCT Ala	ATA Ile	AAT Asn	ATC Ile	GCT Ala 225	AGA Arg	GAT Asp	GCA Ala	AGC Ser	GCT Ala 230	ACA Thr	TTA Leu	TTA Leu	2004
TCC Ser	ACT Thr 235	TGC Cys	TAT Tyr	TAT Tyr	CGC Arg	AAT Asn 240	TGC Cys	CGC Arg	GAG Glu	AGG <b>A</b> rg	ACT Thr 245	ATA Ile	CTT Leu	CGC Arg	CCT Pro	2052
TCT Ser 250	CGT Arg	CTC Leu	GGA Gly	CAT His	GGT Gly 255	TAC Tyr	ACA Thr	ATC Ile	CCT Pro	TCT Ser 260	CCC Pro	GGT Gly	GCC Ala	GAT Asp	ATG Met 265	2100
CTT Leu	TAT Tyr	GAA Glu	GAA Glu	GAC Asp 270	GTA Val	TAT Tyr	AGT Ser	TTT Phe	GAC Asp 275	GCA Ala	GCT Ala	AAA Lys	GGC Gly	CAT His 280	TAT Tyr	2148
TCG Ser	TCA Ser	ATA Ile	TTT Phe 285	CTA Leu	TGT Cys	TAT Tyr	GCC Ala	ATG Met 290	GGG Gly	CTT Leu	ACA Thr	ACA Thr	CCG Pro 295	CTG Leu	ATT Ile	2196
ATT Ile	GCG Ala	CTC Leu 300	CAT His	AAA Lys	TAT Tyr	ATG Met	GCG Ala 305	GGC Gly	ATT Ile	AAA Lys	AAT Asn	TCG Ser 310	TCA Ser	GAT Asp	TGG Trp	2244

Thr	GCT Ala 315	ACA Thr	TTA Leu	CAA Gln	GGC Gly	ATG Met 320	TAC Tyr	GGG Gly	CTT Leu	GTC Val	TTG Leu 325	GGA Gly	TCG Ser	CTA Leu	TCG Ser	2292
TCA Ser 330	Leu	TGT Cys	ATT Ile	CCA Pro	Ser 335	AGC Ser	AAC Asn	AAC Asn	GAT Asp	GCC Ala 340	CTA Leu	ATT Ile	CGT Arg	CCC Pro	ATT Ile 345	2340
CAA Gln	ATT Ile	TTG Leu	ATA Ile	TTG Leu 350	ATA Ile	ATC Ile	GGT Gly	GCA Ala	CTG Leu 355	GCC Ala	ATT Ile	GCA Ala	TTG Leu	GCT Ala 360	GGA Gly	2388
			ATT Ile 365													2436
			TTT Phe													2484
			GCA Ala													2532
			ATG Met										TGA	raga:	rcg	2581
TCG	STCT	GCG (	CATCO	SCCC#	AT GO	CTGG	CGGA	A CGC	CTCT?	TTCG	AAC	GTG	AAT A	AAAA	CTTTGT	2641
ATC'	ract:	AAA	CAATA	ACTI	rt Gi	rgtt7	TAT	GAC	GCGG1	rcga	AAA	CAATO	GAG (	GAGC	rgcaat	2701
ATT	AAGC:	raa (	CCGC	ATACO	3C . CC	GGCC	GGT	AA A	BACCA	ATTT	TATA	ACCA	TAT	racgo	CATCTA	2761
TCG	AAAC'	ITG :	TTCG1	AGAA	CC GC	CAAGT				rcc A Ser A						2813
CGC	GTA	CTG	CGC Arg	CTG	ACG	GGA	TGG	1et \ 1 GTG	/al s GGC	Ser A	Asn M	1et A 5 CTA	rg \ GTT	/al I CTG	Leu TCT	2813
CGC Arg	GTA Val 10 CAG	CTG Leu CAA Gln	CGC	CTG Leu TCT Ser	ACG Thr TGT Cys	GGA Gly 15 GCC Ala	TGG Trp GGA Gly	Met V 1 GTG Val TTG Leu	GGC Gly CCC Pro	ATA Ile CAT	TTT Phe 20 AAC Asn	OTA Leu GTC Val	GTT Val GAT Asp	/al I CTG Leu ACC Thr	TCT Ser CAT His	
CGC Arg TTA Leu 25	GTA Val 10 CAG Gln	CTG Leu CAA Gln	CGC Arg ACC Thr	CTG Leu TCT Ser	ACG Thr TGT Cys 30	GGA Gly 15 GCC Ala	TGG Trp GGA Gly	Met V  1  GTG  Val  TTG  Leu  CCC	GGC Gly CCC Pro	ATA Ile CAT His 35	TTT Phe 20 AAC Asn	Met A 5 CTA Leu GTC Val	GTT Val GAT Asp	/al I CTG Leu ACC Thr	TCT Ser CAT His 40	2861
CGC Arg TTA Leu 25 CAT His	GTA Val 10 CAG Gln ATC Ile	CTG Leu CAA Gln CTA Leu	CGC Arg ACC Thr	CTG Leu TCT Ser TTC Phe 45	ACG Thr TGT Cys 30 AAC Asn	GGA Gly 15 GCC Ala CCT Pro	TGG Trp GGA Gly TCT Ser	1 GTG Val TTG Leu CCC Pro	GGC Gly CCC Pro ATT Ile 50	ATA Ile CAT His 35 TCG Ser	TTT Phe 20 AAC Asn GCC Ala	Met A 5 CTA Leu GTC Val GAT Asp	GTT Val GAT Asp GGC Gly	/al I CTG Leu ACC Thr GTT Val 55	TCT Ser CAT His 40 CCT Pro	2861 2909
CGC Arg TTA Leu 25 CAT His TTG Leu GCC	GTA Val 10 CAG Gln ATC Ile TCA Ser	CTG Leu CAA Gln CTA Leu GAG Glu	CGC Arg ACC Thr ACT Thr GTG Val	CTG Leu TCT Ser TTC Phe 45 CCC Pro	ACG Thr TGT Cys 30 AAC Asn AAT ASn	GGA Gly 15 GCC Ala CCT Pro TCG Ser	TGG Trp GGA Gly TCT Ser CCT Pro	Met V  1  GTG  Val  TTG  Leu  CCC  Pro  ACG  Thr  65	GGC Gly CCC Pro ATT Ile 50 ACC Thr	ATA Ile CAT His 35 TCG Ser GAA Glu	TTT Phe 20 AAC Asn GCC Ala TTA Leu ACT	Met A 5 CTA Leu GTC Val GAT Asp TCT Ser AGT	GTT Val GAT Asp GGC Gly ACA Thr 70	/al I CTG Leu ACC Thr GTT Val 55 ACT Thr	TCT Ser CAT His 40 CCT Pro GTC Val	2861 2909 2957
CGC Arg TTA Leu 25 CAT His TTG Leu GCC Ala	GTA Val 10 CAG Gln ATC Ile TCA Ser ACC Thr	CTG Leu CAA Gln CTA Leu GAG Glu AAG Lys 75	CGC Arg ACC Thr ACT Thr GTG Val 60 ACA	CTG Leu TCT Ser TTC Phe 45 CCC Pro	ACG Thr TGT Cys 30 AAC Asn AAT Asn GTA Val	GGA Gly 15 GCC Ala CCT Pro TCG Ser CCG Pro	TGG Trp GGA Gly TCT Ser CCT Pro ACG Thr 80	Met V  1  GTG Val  TTG Leu  CCC Pro  ACG Thr 65  ACT Thr	GGC Gly CCC Pro ATT Ile 50 ACC Thr GAA Glu	ATA Ile CAT His 35 TCG Ser GAA Glu AGC Ser	TTT Phe 20 AAC Asn GCC Ala TTA Leu ACT Thr	Met A 5 CTA Leu GTC Val GAT Asp TCT Ser AGT Ser 85	GTT Val GAT Asp GGC Gly ACA Thr 70 TCC Ser	/al I CTG Leu ACC Thr GTT Val 55 ACT Thr TCC Ser	TCT Ser CAT His 40 CCT Pro GTC Val	2861 2909 2957 3005

												AAC Asn				3197	
GAC Asp	GTC Val	ACG Thr	TTC Phe 140	AAT Asn	CCA Pro	ATC Ile	GAA Glu	TAC Tyr 145	CAC His	GCC Ala	AAC Asn	GAA Glu	AAG Lys 150	AAT Asn	GTA Val	3245	
GAG Glu	GTT Val	GCC Ala 155	CGA Arg	GTG Val	GCC Ala	GGT Gly	CTA Leu 160	TAC Tyr	GGA Gly	GTA Val	CCG Pro	GGG Gly 165	TCG Ser	GAT Asp	TAT Tyr	3293	
GCA Ala	TAC Tyr 170	CCT Pro	AGG Arg	AAA Lys	TCG Ser	GAA Glu 175	TTA Leu	ATA Ile	TCC Ser	TCC Ser	ATT Ile 180	CGA Arg	CGG Arg	GAT Asp	CCC Pro	3341	
CAG Gln 185	GGT Gly	TCT Ser	TTC Phe	TGG Trp	ACT Thr 190	AGT Ser	CCT Pro	ACA Thr	CCC Pro	CGT Arg 195	GGA Gly	AAT Asn	AAA Lys	TAT Tyr	TTC Phe 200	3389	
ATA Ile	TGG Trp	ATT Ile	AAT Asn	AAA Lys 205	ACA Thr	ATG Met	CAC His	ACC Thr	ATG Met 210	GGC Gly	GTG Val	GAA Glu	GTT Val	AGA Arg 215	AAT Asn	3437	
GTC Val	GAC Asp	TAC Tyr	AAA Lys 220	GAC Asp	AAC Asn	GGC Gly	TAC Tyr	TTT Phe 225	CAA Gln	GTG Val	ATA Ile	CTG Leu	CGT Arg 230	GAT Asp	AGA Arg	3485	
TTT Phe	AAT Asn	CGC Arg 235	CCA Pro	TTG Leu	GTA Val	GAA Glu	AAA Lys 240	CAT His	ATT Ile	TAC Tyr	ATG Met	CGT Arg 245	GTG Val	TGC Cys	CAA Gln	3533	•
CGA Arg	CCC Pro 250	GCA Ala	TCC Ser	GTG Val	GAT Asp	GTA Val 255	TTG Leu	GCC Ala	CCT Pro	CCA Pro	GTT Val 260	CTC Leu	AGC Ser	GGA Gly	GAA Glu	3581	
AAC Asn 265	TAC Tyr	AAA Lys	GCA Ala	TCT Ser	TGC Cys 270	ATC Ile	GTT Val	AGA Arg	CAT His	TTT Phe 275	TAT Tyr	CCC Pro	CCG Pro	GGA Gly	TCT Ser 280	3629	
GTC Val	TAC Tyr	GTA Val	TCT Ser	TGG Trp 285	AGA Arg	CGT Arg	AAC Asn	GGA Gly	AAC Asn 290	ATT Ile	GCC Ala	ACA Thr	CCC Pro	CGC Arg 295	AAG Lys	3677	
GAC Asp	CGT Arg	GAC Asp	GGG Gly 300	AGT Ser	TTT Phe	TGG Trp	TGG Trp	TTC Phe 305	GAA Glu	TCT Ser	GGC Gly	CGC Arg	GGG Gly 310	AIG	ACA Thr	3725	
CTA Leu	GTA Val	TCC Ser 315	Thr	ATA Ile	ACC Thr	CTC Leu	GGA Gly 320	Asn	TCT Ser	GGA Gly	CTC Leu	GAA Glu 325	TCT Ser	CCT Pro	CCA Pro	3773	
AAG Lys	GTT Val 330	Ser	TGC Cys	TTG Leu	GTA Val	GCG Ala 335	Trp	AGG Arg	CAA Gln	GGC Gly	GAT Asp 340	Mer	ATA Ile	AGC Ser	ACA Thr	3821	
TCG Ser 345	AAT Asn	GCT Ala	ACA Thr	GCT Ala	GTA Val 350	Pro	ACG Thr	GTA Val	TAT Tyr	TAT Tyr 355	HIS	CCC Pro	CGT Arg	ATC	TCT Ser 360	3869	
CTG Leu	GCA Ala	TTT Phe	AAA Lys	GAT Asp 365	Gly	TAT Tyr	GCA Ala	ATA Ile	TGT Cys 370	Thr	ATA	GAA Glu	TGI	GTI Val	CCC	3917	
TCT Ser	GGG Gly	ATT Ile	ACT Thr 380	Val	AGG Arg	TGG Trp	TTA Leu	GTI Val 385	His	GAT Asp	GAA Glu	CCC Pro	CAG Glr 390	PIC	AAC Asn	3965	

ACA Thr	3 00															
	Thr	TAT Tyr 395		ACT Thr	GTG Val	GTT Val	ACA Thr 400	Gly	CTC Leu	TGC Cys	AGG Arg	ACC Thr 405	ATC Ile	GAT Asp	CGT Arg	4013
TAT Tyr	AGA Arg 410	Asn	CTC Leu	GCC Ala	AGT Ser	CGG Arg 415	ATT Ile	CCA Pro	GTC Val	CAG Gln	GAC Asp 420	AAC Asn	TGG Trp	GCG Ala	AAA Lys	4061
ACG Thr 425	Lys	TAT Tyr	ACG Thr	TGC Cys	AGA Arg 430	CTA Leu	ATT Ile	GGA Gly	TAT Tyr	CCG Pro 435	TTC Phe	GAC Asp	GTG Val	GAT Asp	AGA Arg 440	4109
			TCC Ser													4157
															TTA Leu	4205
GGT Gly	ATT Ile	GGT Gly 475	ATC Ile	ATT Ile	ATC Ile	ACA Thr	GCC Ala 480	CTA Leu	TGC Cys	TTT Phe	TAC Tyr	CTA Leu 485	CCG Pro	GGG Gly	CGG Arg	4253
AAT Asn		GATT	AAC (	CATC	GTAT(	GT GA	TATA	AAAA	A TT	ATTA	AGTG	TTA'	T <b>AAC</b> (	CGA		4306
TCG	CATT	CTT (	CTG <b>T</b> I	rtcg/	T TA	CACA	AATA	A TA	<b>LAA</b> F	GGTA	TTG	TAAT	CAG	CACC	ATCGCA	4366
TTG	TTTC	GTA (	GATG!	ACTC	AT G	rtca(	STCC	G CG	rga <b>t</b> (	GTCA	LAAA	ATAC	GTA :	TTTT?	rggtat	4426
CAC	GCAG(	CGG (	CAA	AATG	cc cz	ATTA	rgtti	A TT	ומדדו	ידרר	אממ	2000	י מיחיר	מידירים	AAACAT	4486
										-100	AAA.		31 <b>A</b> .	LILI	Dutchi	1100
CGG	GACG'	rac A	ATÇAT												ATCGC	4546
				rgtgo	GC GC	CACG	CTAA:	r cg:	CATA)	CGGT	GCC	GCTA(	CAT :	TAAA		
AAG	rct <b>c</b> (	CGA 1	ATATO	rgtg( Caag(	GC GC	CACGI	TAA:	r cg:	rata( GTCG(	CGG <b>T</b> STAA	GCC	GCTA(	CAT 1	TAAA CATC	AATCGC	4546
AAG GAT	rctc( ACGG)	CGA 1	ATATO	rgtg( Caag( Acaa)	GC GC	CACGO ACGGO CTGAO	TTAA: CCAA! STAG!	r cg: A Acc	PATAC GTCGC PCCTI GA AT	CGGT ETAA ATAT IG AT	GCCC TAAT AGTT	GCTACTO	CAT TACG C	TAAA CATCO TAGTO	AATCGC GAATGT GATACA	4546 4606
AAG' GAT. CAA' GAA	TCTC(ACGG)	CGA AAAA AAAA A	ATATO	TGTGC CAAGO ACAAT CTGGC GAT	GC GC CT CX CC GC CTC	CACGT ACGGC CTGAC ATATC	CAAA GTAGA CATAC GTG	r cgr A Acc A TTT r AAC	GTCGCTATE  ACCTATE  A	CGGT GTAA ATAT TG AT et Me 1	GCCC TAA: AGTT G TC et Se	GCTAG FCTTA FACTG CG CG PI	CAT :	TAAAA	AATCGC GAATGT GATACA CT CO	4546 4606 4666
GAA GAT CAA GAA Glu	CTCCACCACCACCACCACCACCACCACCACCACCACCACC	CGA ATA (AAAA AGAT ASp	ATATO CCGTA ATCGO CGC Arg	TGTGC CAAGC ACAAT CTGGC GAT Asp	GC GC CT CA CTC Leu ACA	CACGT ACGGC CTGAC ATATC GTT Val	CTAA: CCAA; CATAC GTG Val	T CGT A ACC A TTT T AAC GTT Val 15 GAG	TATAGETCGG TCCTA GA AT Me CGT Arg	GGGT  ATAT  TG AT  et Me  1  GGA  Gly  CGA	GCCC TAA: AGT: G TC et Se CGT Arg	CTC CTC CTC CTC CTC Leu	CAT TACG OF CAC TO THE CAC Arg 20 CGT	TAAAAATCATCATCATCATCATCATCATCATCATCATCAT	AATCGC GAATGT GATACA CT CO ATG Met	4546 4606 4666 4718
GAA GAA GAA Glu GAT Asp	CGA	GAT ASP GGC Gly 25	ATATO CCGTA ATCGO CGC Arg 10 ACG	CAAGO ACAAT CTGGO GAT Asp GAA Glu TGT	GC GC CT CA CC GC GG TA CTC Leu ACA Thr	CACGT ACGGC ATATO GTT Val GAT Asp	CTAA: CCAA; CATAC CATAC GTG Val AGA Arg 30	T CGT A ACC A TTT T AAC GTT Val 15 GAG Glu ACG	TATAGET CAA GIN	GGGT GTAA ATAT TG AT et Me 1 GGA Gly CGA Arg	GCCC TAA: AGT: GCCET Set Se CGT Arg CAT His	CTC Leu  CCA Pro 35	CAT : ACG C CAG T CC AC TO Th CGA Arg 20 CGT Arg	TAAAA CATCO TAGTO CC CO nr Pr 5 ATG Met ACG Thr	AATCGC GAATGT GATACA CT CO ATG Met ACT Thr	4546 4606 4666 4718 4766
GAA Glu GAT Asp	CGA Arg	GAT ASP GGC Gly 25 TCG Ser CTC	CGC Arg 10 ACG Thr	GAAGO GAT ASP GAA Glu TGT Cys	CTC GO CTC Leu ACA Thr TGT Cys	CACGO ACGGO GTT Val GAT Asp GGG GGY 45	CCAAA GTAGA CATAC GTG Val AGA AGA AGA TGT Cys	A ACC A TTT T AAC GTT Val 15 GAG Glu ACG Thr	CGT Arg CAA Gln ATA Ile	GGA	GCCC TAAT AGTT GG TG et Se CGT Arg CAT His	CTC CCA Pro 35 GTA Val	CAT TAGE OF THE CGA Arg 20 CGT Arg TTT Phe	TAAAA CATCO TAGTO CC CO TAGTO S ATG Met ACG Thr ACC Thr	AATCGC GAATGT GATACA CT CO ATG Met ACT Thr ATA Ile	4546 4606 4666 4718 4766
GAAGIU GATASP TGGTrp TTC Phe 55	GAT ASP AGC Ser CGA Arg 40 GTT Val	GAT ASP GGC Gly 25 TCG Ser CTC Leu GCC	CGC Arg 10 ACG Thr	GAAGO GAT Asp GAA Glu TGT Cys	CTC Leu ACA Thr Cys GCA Ala 60	CACGT ACGGG ATATO  GTT Val  GAT Asp  GGG Gly 45 GTA Val	CCAAA GTAGA GTG Val AGA AGA TGT Cys	A ACCA TTT AACCA GTT Val 15 GAG Glu ACG Thr TTG Leu	CGT ATA Gln  ATA Gly  CCC	GGA	GCCC TAAT AGTT GGTG et Se CGT Arg CAT His ATG Met 50 CTA Leu	CTC CCA Pro 35 GTA Val	CAT : ACG (CAC) CC AC CC AC CC AC CC AC CC AC Th  CGA Arg 20 CGT Arg TTT Phe ACT Thr	TAAAA CATCO TAGTO CC CO TAGTO S ATG Met ACG Thr ACC Thr GTT Val	AATCGC GAATGT GATACA CT CO ATG Met ACT Thr ATA Ile TCA Ser 70 TTG	4546 4606 4666 4718 4766 4814

GCG Ala	TTG Leu	GAT Asp 105	ACA Thr	TGT Cys	GCT Ala	CGG Arg	CAT His 110	AAC Asn	AGC Ser	AAA Lys	CTT Leu	ATT Ile 115	GAC Asp	TTC Phe	GCA Ala	5054
AAC Asn	GCC Ala 120	AAA Lys	GTT Val	CTG Leu	GTT Val	GAA Glu 125	GCT Ala	ATC Ile	GCC Ala	CCA Pro	TTC Phe 130	GGT Gly	GTG Val	CCA Pro	AAT Asn	5102
GCA Ala 135	GCA Ala	TAT Tyr	GGG Gly	GAA Glu	GTC Val 140	TTC Phe	CGG Arg	TTA Leu	AGG Arg	GAC Asp 145	AGC Ser	AAA Lys	ACC Thr	ACG Thr	TGT Cys 150	5150
		CCT Pro														5198
		GTT Val														5246
		AGA Arg 185														5294
		GTC Val												TAAZ	<b>AAC</b> GCA	5346
CCTC	TAAC	GG I	TACI	GTGT	T TA	TTAT	CCAF	A TCA	CAC	CATA	GACA	TTAT	TA C	TAAT	ATATG	5406
GATO	TTTA	TT I	CATA	TAAT	r <b>G</b>											5426

### (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 369 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Met Ala Gly Ile Thr Val Ala Cys Asp His Thr Ala Gly Glu Ala

His Thr Pro Glu Asp Met Gln Lys Lys Trp Arg Ile Ile Leu Ala Gly 20 25 30

Glu Lys Phe Met Thr Ile Ser Ala Ser Leu Lys Ser Ile Val Ser Cys 35 40 45

Val Lys Asn Pro Leu Leu Thr Phe Gly Ala Asp Gly Leu Ile Val Gln
50 60

Gly Thr Val Cys Gly Gln Arg Ile Phe Val Pro Ile Asp Arg Asp Ser
65 70 75 80

Phe Ser Glu Tyr Glu Trp His Gly Pro Thr Ala Met Phe Leu Ala Leu 85 90 95

Thr Asp Ser Arg Arg Thr Leu Leu Asp Ala Phe Lys Cys Glu Lys Arg

Arg Ala Ile Asp Val Ser Phe Thr Phe Ala Gly Glu Pro Pro Cys Arg

His Leu Ile Gln Ala Val Thr Tyr Met Thr Asp Gly Gly Ser Val Ser 135 Asn Thr Ile Ile Lys Tyr Glu Leu Trp Asn Ala Ser Thr Ile Phe Pro 150 155 Gln Lys Thr Pro Asp Val Thr Phe Ser Leu Asn Lys Gln Gln Leu Asn Lys Ile Leu Ala Val Ala Ser Lys Leu Gln His Glu Glu Leu Val Phe Ser Leu Lys Pro Glu Gly Gly Phe Tyr Val Gly Thr Val Cys Thr Val 200 Ile Ser Phe Glu Val Asp Gly Thr Ala Met Thr Gln Tyr Pro Tyr Asn Pro Pro Thr Ser Ala Thr Leu Ala Leu Val Val Ala Cys Arg Lys 235 Lys Ala Asn Lys Asn Thr Ile Leu Thr Ala Tyr Gly Ser Gly Lys Pro Phe Cys Val Ala Leu Glu Asp Thr Ser Ala Phe Arg Asn Ile Val Asn 265 Lys Ile Lys Ala Gly Thr Ser Gly Val Asp Leu Gly Phe Tyr Thr Thr Cys Asp Pro Pro Met Leu Cys Ile Arg Pro His Ala Phe Gly Ser Pro 295 Thr Ala Phe Leu Phe Cys Asn Thr Asp Cys Met Thr Ile Tyr Glu Leu

Glu Glu Val Ser Ala Val Asp Gly Ala Ile Arg Ala Lys Arg Ile Asn

Glu Tyr Phe Pro Thr Val Ser Gln Ala Thr Ser Lys Lys Arg Lys Gln

Ser Pro Pro Pro Ile Glu Arg Glu Arg Lys Thr Thr Arg Ala Asp Thr

Gln

## (2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 422 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Pro Ser Gly Ala Ser Ser Pro Pro Pro Ala Tyr Thr Ser Ala

Ala Pro Leu Glu Thr Tyr Asn Ser Trp Leu Ser Ala Phe Ser Cys Ala 25

Tyr	Pro	Gln 35	Cys	Thr	Ala	Gly	Arg 40	Gly	His	Arg	Gln	Asn 45	G <u>l</u> y	Lys	Lys
Cys	Ile 50	Arg	Cys	Ile	Val	Ile 55	Ser	Val	Cys	Ser	Leu 60	Val	Cys	Ile	Ala
Ala 65	His	Leu	Ala	Val	Thr 70	Val	Ser	Gly	۷al	Ala 75	Leu	Ile	Pro	Leu	Ile 80
Asp	Gln	Asn	Arg	Ala 85	Tyr	Gly	Asn	Cys	Thr 90	Val	Cys	Val	Ile	Ala 95	Gly
Phe	Ile	Ala	Thr 100	Phe	Ala	Ala	Arg	Leu 105	Thr	Ile	Arg	Leu	Ser 110	Glu	Thr
Leu	Met	Leu 115	Val	Gly	Lys	Pro	Ala 120	Gln	Phe	Ile	Phe	Ala 125	Ile	Ile	Ala
Ser	Val 130		Glu	Thr	Leu	Ile 135		Asn	Glu	Ala	Leu 140		Ile	Ser	Asn
Thr 145	Thr	Tyr	Lys	Thr	Ala 150	Leu	Arg	Ile	Ile	Glu 155	Val	Thr	Ser	Leu	Ala 160
Cys	Phe	Val	Met	Leu 165	Gly	Ala	Ile	Ile	Thr 170	Ser	His	Asn	Tyr	Val 175	Cys
Ile	Ser	Thr	Ala 180	Gly	Asp	Leu	Thr	Trp 185	Lys	Gly	Gly	Ile	Phe 190	His.	Ala
Tyr	His	Gly 195	Thr	Leu	Leu	Gly	Ile 200	Thr	Ile	Pro	Asn	Ile 205	His	Pro	Ile
Pro	Leu 210	Ala	Gly	Phe	Leu	Ala 215	Val	Tyr	Thr	Ile	Leu 220	Ala	Ile	Asn	Ile
Ala 225	Arg	Asp	Ala	Ser	Ala 230	Thr	Leu	Leu	Ser	Thr 235	Cys	Tyr	Tyr	Arg	Asn 240
Cys	Arg	Glu	Arg	Thr 245	Ile	Leu	Arg	Pro	Ser 250	Arg	Leu	Gly	His	Gly 255	Tyr
Thr	Ile	Pro	Ser 260	Pro	Gly	Ala	Asp	Met 265	Leu	Tyr	Glu	Glu	Asp 270	Val	Tyr
Ser	Phe	Asp 275	Ala	Ala	Lys	Gly	His 280	Tyr	Ser	Ser	Ile	Phe 285	Leu	Cys	Tyr
Ala	Met 290	Gly	Leu	Thr	Thr	Pro 295	Leu	Ile	Ile	Ala	Leu 300	His	Lys	Tyr	Met
Ala 305	Gly	Ile	Lys	Asn	Ser 310	Ser	Asp	Trp	Thr	Ala 315	Thr	Leu	Gln	Gly	Met 320
Tyr	Gly	Leu	Val	Leu 325	Gly	Ser	Leu	Ser	Ser 330	Leu	Cys	Ile	Pro	Ser 335	Ser
Asn	Asn	Asp	Ala 340	Leu	Ile	Arg	Pro	Ile 345	Gln	Ile	Leu	Ile	Leu 350	Ile	Ile
Gly	Ala	Leu 355	Ala	Ile	Ala	Leu	Ala 360	Gly	Cys	Gly	Gln	Ile 365	Ile	Gly	Pro
Thr	Leu 370	Phe	Ala	Ala	Ser	Ser 375	Ala	Ala	Met	Ser	Cys 380	Phe	Thr	Cys	Ile

Asn Ile Arg Ala Thr Asn Lys Gly Val Asn Lys Leu Ala Ala Ala Ser 385 390 395

Val Val Lys Ser Val Leu Gly Phe Ile Ile Ser Gly Met Leu Thr Cys 405 410 415

Val Leu Leu Pro Leu Ser 420

#### (2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 489 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Val Ser Asn Met Arg Val Leu Arg Val Leu Arg Leu Thr Gly Trp

1 10 15

Val Gly Ile Phe Leu Val Leu Ser Leu Gln Gln Thr Ser Cys Ala Gly
20 25 30

Leu Pro His Asn Val Asp Thr His His Ile Leu Thr Phe Asn Pro Ser

Pro Ile Ser Ala Asp Gly Val Pro Leu Ser Glu Val Pro Asn Ser Pro 50 55 60

Thr Thr Glu Leu Ser Thr Thr Val Ala Thr Lys Thr Ala Val Pro Thr 65 70 75 80

Thr Glu Ser Thr Ser Ser Ser Glu Ala His Arg Asn Ser Ser His Lys
85 90 95

Ile Pro Asp Ile Ile Cys Asp Arg Glu Glu Val Phe Val Phe Leu Asn 100 105 110

Asn Thr Gly Arg Ile Leu Cys Asp Leu Ile Val Asp Pro Pro Ser Asp 115 120 125

Asp Glu Trp Ser Asn Phe Ala Leu Asp Val Thr Phe Asn Pro Ile Glu 130 135 140

Tyr His Ala Asn Glu Lys Asn Val Glu Val Ala Arg Val Ala Gly Leu 145 150 155 160

Tyr Gly Val Pro Gly Ser Asp Tyr Ala Tyr Pro Arg Lys Ser Glu Leu 165 170 175

Ile Ser Ser Ile Arg Arg Asp Pro Gln Gly Ser Phe Trp Thr Ser Pro 180 185 190

Thr Pro Arg Gly Asn Lys Tyr Phe Ile Trp Ile Asn Lys Thr Met His 195 200 205

Thr Met Gly Val Glu Val Arg Asn Val Asp Tyr Lys Asp Asn Gly Tyr 210 220

Phe Gln Val Ile Leu Arg Asp Arg Phe Asn Arg Pro Leu Val Glu Lys 225 230 235

167

His Ile Tyr Met Arg Val Cys Gln Arg Pro Ala Ser Val Asp Val Leu Ala Pro Pro Val Leu Ser Gly Glu Asn Tyr Lys Ala Ser Cys Ile Val Arg His Phe Tyr Pro Pro Gly Ser Val Tyr Val Ser Trp Arg Arg Asn 280 Gly Asn Ile Ala Thr Pro Arg Lys Asp Arg Asp Gly Ser Phe Trp Trp Phe Glu Ser Gly Arg Gly Ala Thr Leu Val Ser Thr Ile Thr Leu Gly Asn Ser Gly Leu Glu Ser Pro Pro Lys Val Ser Cys Leu Val Ala Trp 330 Arg Gln Gly Asp Met Ile Ser Thr Ser Asn Ala Thr Ala Val Pro Thr Val Tyr Tyr His Pro Arg Ile Ser Leu Ala Phe Lys Asp Gly Tyr Ala Ile Cys Thr Ile Glu Cys Val Pro Ser Gly Ile Thr Val Arg Trp Leu Val His Asp Glu Pro Gln Pro Asn Thr Thr Tyr Asp Thr Val Val Thr 395 390 Gly Leu Cys Arg Thr Ile Asp Arg Tyr Arg Asn Leu Ala Ser Arg Ile Pro Val Gln Asp Asn Trp Ala Lys Thr Lys Tyr Thr Cys Arg Leu Ile Gly Tyr Pro Phe Asp Val Asp Arg Phe Gln Asn Ser Glu Tyr Tyr Asp Ala Thr Pro Ser Ala Arg Gly Met Pro Met Ile Val Thr Ile Thr Ala 455 Val Leu Gly Leu Ala Leu Phe Leu Gly Ile Gly Ile Ile Thr Ala Leu Cys Phe Tyr Leu Pro Gly Arg Asn 485

## (2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 212 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met Met Ser Pro Thr Pro Glu Asp Asp Arg Asp Leu Val Val Arg
1 5 10 15

Gly Arg Leu Arg Met Met Asp Ser Gly Thr Glu Thr Asp Arg Glu Gln 20 25 30

**** 9	His	Pro 35	Arg	Thr	Thr	Trp	Arg 40	Ser	Ile	Cys	Cys	_Gly 45	Cys	Thr	Ile	
Gly	Met 50	Val	Phe	Thr	Ile	Phe 55	Val	Leu	Val	Ala	Ala 60	Val	Leu	Leu	Gly	
Ser 65	Leu	Phe	Thr	Val	Ser 70	Tyr	Met	Ala	Met	Glu 75	Ser	Gly	Thr	Cys	Pro 80	
Asp	Glu	Trp	Ile	Gly 85	Leu	Gly	Tyr	Ser	Cys 90	Met	Arg	Val	Ala	Gly 95	Lys	
Asn	Ala	Thr	Asp 100	Leu	Glu	Ala	Leu	Asp 105	Thr	Cys	Ala	Arg	His 110	Asn	Ser	
Lys	Leu	Ile 115	Asp	Phe	Ala	Asn	Ala 120	Lys	Val	Leu	Val	Glu 125	Ala	Ile	Ala	
Pro	Phe 130	Gly	Val	Pro	Asn	Ala 135	Ala	Tyr	Gly	Glu	Val 140	Phe	Arg	Leu	Arg	
Asp 145	Ser	Lys	Thr	Thr	Cys 150	Ile	Arg	Pro	Thr	Met 155	Gly	Gly	Pro	Val	Ser 160	
Ala	Asp	Cys	Pro	Val 165	Thr	Cys	Thr	Val	Ile 170	Cys	Gln	Arg	Pro	Arg 175	Pro	
Leu	Ser	Thr	Met 180	Ser	Ser	Ile	Ile	Arg 185	Asp	Alď	Arg	Val	Tyr 190	Leu	His	
Leu	Glu	Arg 195	Arg	Asp	Tyr	Tyr	Glu 200	Val	Tyr	Ala	Ser	Val 205	Leu	Ser	Asn	
Ala	Met 210	Ser	Lys													
(2)	INFO	ORMA:	rion	FOR	SEQ	ID 1	10 : 8 :	:								
	(i)	( )	QUENC		IARAC I: 15											
		((	3) TY 2) ST 2) TO	PE:	EDNE	ESS:	acid doub	ì	rs							
	(ii)	(1	3) TY	(PE: TRANI OPOLO	EDNE GY:	ESS: line	acid doub ear	l ole								
	(ii) (iii)	O) IOM	B) TY C) ST C) TO LECUI	(PE: TRANI OPOLO LE TY	EDNE GY: PE:	ESS: line DNA	acid doub ear	l ole								
	(iii) (iv)	1) 10M 10H 1YH	3) TY C) ST C) TO LECUI POTHE	(PE: TRANI )POLO LE TY ETICA ENSE:	DEDNE DGY: PE:	ESS: line DNA	acid doub ear	l ole								
	(iii) (iv)	MOI HYI ANT	B) TY C) ST C) TO LECUI	(PE: TRANI OPOLO LE TY ETICA CNSE: L: LME/K	DEDNIEDGY:  (PE: AL: NO  NO	ESS: line DNA NO	acid doub ear (ger	l ole								
	(iii) (iv) (ix)	MOI HYI ANT	3) TY C) ST C) TO LECUI POTHE FI-SE ATURE A) NE	(PE: TRANI DPOLO LE TY TTICA CNSE: L: LME/KOCATI	DEDNIE DGY:  PE:  NO  KEY:  ON:	ESS: line DNA NO CDS	acid douk ear (ger	i ole nomic	:)	):8:						
ATG	(iii) (iv) (ix)	MOI HYI ANT FEA (I SEC	3) TY C) ST C) TC ECUI POTHE FI-SE ATURE ATURE A) NA B) LC QUENC	PE: TRANI POLC  ETICA  ETICA  CNSE: ME/K CATI  CE DE	DEDNE DGY: PE: AL: NO NO CEY: CON: CSCRI	ESS: line DNA NO CDS 11	acidoulear (ger .506 ON: S	i ole nomic SEQ I	D NC	GGG						48
ATG Met 1	(iii) (iv) (ix) (xi)	MOI HYI ANT FEA (I SEC ACG Thr	3) TY C) ST C) TC CECUI POTHE TI-SE ATURE A) NA B) LC CCG Pro TTA	CE TY CASE: CASE: CATICA CATIC	DEDNE DGY: TPE: AL: NO EY: CON: CSCRI GTG Val	ESS: line DNA NO CDS 11 PTIC TTA Leu	acid doub ear (ger 506 ON: S CGA Arg	i ole nomic SEQ I GCT Ala	TTG Leu 10	GGG Gly GGA	Trp	Thr	Gly CTT	Leu 15 AGC	Phe	48

		35					40					45				
													TCC Ser			192
													ATA Ile			240
													ACG Thr			288
													TGC Cys 110			336
													GTT Val			384
													TTT Phe			432
GAT Asp 145	Leu	ATC Ile	TTT Phe	AAC Asn	CCA Pro 150	Ile	GAA Glu	TAC Tyr	CAC His	GCC Ala 15	Asn	GAA Glu	AAG Lys	AAT Asn	GTG Val 160	480
													TCA Ser			528
GCA Ala	TAC Tyr	CCA Pro	CGT Arg 180	CAA Gln	TCT Ser	GAA Glu	TTA Leu	ATT Ile 185	TCT Ser	TCG Ser	ATT Ile	C <b>GA</b> Arg	CGA Arg 190	GAT Asp	CCC Pro	576
CAG Gln	GGC <b>G</b> ly	ACA Thr 195	TTT Phe	TGG Trp	ACG Thr	AGC Ser	CCA Pro 200	TCA Ser	CCT Pro	CAT His	GGA Gly	AAC Asn 205	AAG Lys	TAC Tyr	TTC Phe	624
ATA Ile	TGG Trp 210	ATA Ile	AAC Asn	AAA Lys	ACA Thr	ACC Thr 215	AAT Asn	ACG Thr	ATG Met	GGC Gly	GTG Val 220	G <b>AA</b> Glu	ATT Ile	AGA Arg	AAT Asn	672
GTA Val 225	GAT Asp	TAT Tyr	GCT Ala	GAT Asp	AAT Asn 230	GGC Gly	TAC Tyr	ATG Met	CAA Gln	GTC Val 235	ATT Ile	ATG Met	CGT Arg	GAC Asp	CAT His 240	720
Phe	Asn	Arg	Pro	Leu 245	Ile	Asp	Lys	His	11e 250	Tyr	Ile	Arg	GTG Val	Cys 255	Gln	768
CGA Arg	CCT Pro	GCA Ala	TCA Ser 260	GTG Val	GAT Asp	GTA Val	CTG <b>Le</b> u	GCC Ala 265	CCT Pro	CCA Pro	GTC Val	CTC Leu	AGC Ser 270	GGA Gly	G <b>AA</b> Glu	816
Asn	Tyr	Lys 275	Ala	Ser	Cys	Ile	Val 280	Arg	His	Phe	Tyr	Pro 285	Pro	Gly		864
GTC Val	TAT Tyr 290	GTA Val	TCT Ser	TGG Trp	AGA Arg	CAG Gln 295	AAT Asn	GGA Gly	AAC Asn	ATT Ile	GCA Ala 300	ACT Thr	CCT Pro	CGG Arg	AAA Lys	912
GAT	CGC	GAT	GGA	AGT	TTT	TGG	TGG	TTC	GAA	TCT	GGT	AGA	GGA	GCT	ACG	960

Asp 305	Arg	Asp	Gly	Ser	Phe 310	Trp	Trp	Phe	Glu	Ser 315	Gly	Arg	Gly	Ala	Thr 320	
TTG Leu	GTT Val	TCT Ser	ACA Thr	ATA Ile 325	ACA Thr	TTG Leu	GGA Gly	AAT Asn	TCA Ser 330	GGA Gly	ATT Ile	GAT Asp	TTC Phe	CCC Pro 335	CCC Pro	100
AAA Lys	ATA Ile	TCT Ser	TGT Cys 340	CTG Leu	GTT Val	GCC Ala	TGG Trp	AAG Lys 345	CAG Gln	GGT Gly	GAT Asp	ATG Met	ATC Ile 350	AGC Ser	ACG Thr	105
ACG Thr	AAT Asn	GCC Ala 355	ACA Thr	GCT Ala	ATC Ile	CCG Pro	ACG Thr 360	GTA Val	TAT Tyr	CAT His	CAT His	CCC Pro 365	CGT Arg	TTA Leu	TCC Ser	1104
CTG Leu	GCT Ala 370	TTT Phe	AAA Lys	GAT Asp	GGG Gly	TAT Tyr 375	GCA Ala	ATA Ile	TGT Cys	ACT Thr	ATA Ile 380	GAA Glu	<b>TGT</b> Cys	GTC Val	CCC Pro	115:
TCT Ser 385	GAG Glu	ATT Ile	ACT Thr	GTA Val	CGG Arg 390	TGG Trp	TTA Leu	GTA Val	CAT His	GAT Asp 395	GAA Glu	GCG Ala	CAG Gln	CC <b>T</b> Pro	AAC Asn 400	1200
ACA Thr	ACT Thr	TAT Tyr	AAT Asn	ACT Thr 405	GTG Val	GTT Val	ACA Thr	GGT Gly	CTC Leu 410	TGC Cys	CGG Arg	ACC Thr	ATC Ile	GAT Asp 415	CGC Arg	1248
CAT His	AGA Arg	AAT Asn	CTC Leu 420	CTC Leu	AGC Ser	CGC Arg	ATT Ile	CCA Pro 425	GTA Val	TGG Trp	GAC Asp	AAT Asn	TGG Trp 430	ACG Thr	AAA Lys	1296
ACA Thr	AAA Lys	TAT Tyr 435	ACG Thr	TGC Cys	AGA Arg	CTC Leu	ATA Ile 440	GGC Gly	TAC Tyr	CCC Pro	TTC Phe	GAT Asp 445	GAA Glu	GAT Asp	AAA Lys	1344
TTT Phe	CAA Gln 450	GAT Asp	TCG Ser	GAA Glu	TAT Tyr	TAC Tyr 455	GAT Asp	GCA Ala	ACT Thr	CCA Pro	TCT Ser 460	GCA Ala	AGA Arg	GGA Gly	ACA Thr	1392
CCC Pro 465	ATG Met	G <b>T</b> T Val	ATT Ile	ACG Thr	GTT Val 470	ACG Thr	GCA Ala	GTT Val	TTG Leu	GGA Gly 475	TTG Leu	GCT Ala	GTA Val	ATT Ile	TTA Leu 480	1440
GGG Gly	ATG Met	GGG Gly	ATA Ile	ATC Ile 485	ATG Met	ACT Thr	GCC Ala	CTA Leu	TGT Cys 490	TTA Leu	TAC Tyr	AAC Asn	TCC Ser	ACA Thr 495	CGA Arg	1488
AAA Lys	AAT Asn	ATT Ile	CGA Arg 500	TTA Leu	TAA											1506

# (2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 501 amino acids
  - (B) TYPE: amino acid(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Met Leu Thr Pro Arg Val Leu Arg Ala Leu Gly Trp Thr Gly Leu Phe 1 5 15

Phe Leu Leu Ser Pro Ser Asn Val Leu Gly Ala Ser Leu Ser Arg

20 25 30 Asp Leu Glu Thr Pro Pro Phe Leu Ser Phe Asp Pro Ser Asn Ile Ser Ile Asn Gly Ala Pro Leu Thr Glu Val Pro His Ala Pro Ser Thr Glu 55 Ser Val Ser Thr Asn Ser Glu Ser Thr Asn Glu His Thr Ile Thr Glu Thr Thr Gly Lys Asn Ala Tyr Ile His Asn Asn Ala Ser Thr Asp Lys Gln Asn Ala Asn Asp Thr His Lys Thr Pro Asn Ile Leu Cys Asp Thr 100 105 110 Glu Glu Val Phe Val Phe Leu Asn Glu Thr Gly Arg Phe Val Cys Thr Leu Lys Val Asp Pro Pro Ser Asp Ser Glu Trp Ser Asn Phe Val Leu Asp Leu Ile Phe Asn Pro Ile Glu Tyr His Ala Asn Glu Lys Asn Val 150 Glu Ala Ala Arg Ile Ala Gly Leu Tyr Gly Val Pro Gly Ser Asp Tyr Ala Tyr Pro Arg Gln Ser Glu Leu Ile Ser Ser Ile Arg Arg Asp Pro Gln Gly Thr Phe Trp Thr Ser Pro Ser Pro His Gly Asn Lys Tyr Phe Ile Trp Ile Asn Lys Thr Thr Asn Thr Met Gly Val Glu Ile Arg Asn 215 210 Val Asp Tyr Ala Asp Asn Gly Tyr Met Gln Val Ile Met Arg Asp His 235 Phe Asn Arg Pro Leu Ile Asp Lys His Ile Tyr Ile Arg Val Cys Gln Arg Pro Ala Ser Val Asp Val Leu Ala Pro Pro Val Leu Ser Gly Glu 265 Asn Tyr Lys Ala Ser Cys Ile Val Arg His Phe Tyr Pro Pro Gly Ser Val Tyr Val Ser Trp Arg Gln Asn Gly Asn Ile Ala Thr Pro Arg Lys 295 Asp Arg Asp Gly Ser Phe Trp Trp Phe Glu Ser Gly Arg Gly Ala Thr 310 Leu Val Ser Thr Ile Thr Leu Gly Asn Ser Gly Ile Asp Phe Pro Pro Lys Ile Ser Cys Leu Val Ala Trp Lys Gln Gly Asp Met Ile Ser Thr 345 Thr Asn Ala Thr Ala Ile Pro Thr Val Tyr His His Pro Arg Leu Ser 365 360 355

Leu Ala Phe Lys Asp Gly Tyr Ala Ile Cys Thr Ile Glu Cys Val Pro

	370					375					380					
Ser 385	Glu	Ile	Thr	Val	Arg 390	Trp	Leu	Val	His	Asp 3 <b>9</b> 5	Glu	Ala	Gln	Pro	Asn 400	
Thr	Thr	Tyr	Asn	Thr 405	Val	Val	Thr	Gly	Leu 410	Cys	Arg	Thr	Ile	Asp 415	Arg	
His	Arg	Asn	Leu 420	Leu	Ser	Arg	Ile	Pro 425	Val	Trp	Asp	Asn	Trp 430	Thr	Lys	
Thr	Lys	Tyr 435	Thr	Cys	Arg	Leu	Ile 440	Gly	Tyr	Pro	Phe	Asp 445	Glu	Asp	Lys	
Phe	Gln 450		Ser	Glu	Tyr	Tyr 455	Asp	Ala	Thr	Pro	Ser 460	Ala	Arg	Gly	Thr	
Pro 465	Met	Val	Ile	Thr	Val 470	Thr	Ala	Val	Leu	Gly <b>47</b> 5	Leu	Ala	Val	Ile	Leu 480	
Gly	Met	Gly	Ile	Ile 485	Met	Thr	Ala	Leu	Cys 490	Leu	Tyr	Asn	Ser	Thr 495	Arg	
Lys	Asn	Ile	Arg 500	Leu												
(2)	INF	ORMA!	rion	FOR	SEQ	ID <del>1</del>	10:10	0:								
	(i)	() () ()	A) LI B) T C) S	ENGT YPE : TRAN	HARAC H: 17 nucl DEDNI DGY:	734 l leic ESS:	ase acio doul	pain i	rs							
	(ii)	) MOI	LECUI	LE T	YPE:	DNA	(ger	nomic	2)							
,	(iii)	HYI	РОТНІ	ETIC	AL: 1	10										
	(iv)	ANT	rI-si	ENSE	: NO											
	(ix)		A) NA	AME/I	KEY:		.734									
	(xi)	SEC	QUENC	CE DI	ESCRI	PTIC	N: 5	SEQ 1	D NO	0:10:	:					
ATG Met 1	GAC Asp	CGC Arg	GCC Ala	GTT Val 5	AGC Ser	CAA Gln	GTT Val	GCG Ala	TTA Leu 10	GAG Glu	AAT Asn	GAT Asp	GAA Glu	AGA Arg 15	GAG Glu	48
GCA Ala	AAA Lys	AAT Asn	ACA Thr 20	TGG Trp	CGC <b>A</b> rg	TTG Leu	ATA Ile	TTC Phe 25	CGG Arg	ATT Ile	GCA Ala	ATC Ile	TTA Leu 30	TTC Phe	TTA Leu	96
ACA Thr	GTA Val	GTG Val 35	ACC Thr	TTG Leu	GCT Ala	ATA Ile	TCT Ser 40	GTA Val	GCC Ala	TCC Ser	CTT Leu	TTA Leu 45	TAT Tyr	AGC Ser	ATG Met	144
GGG Gly	GCT Ala 50	AGC Ser	ACA Thr	CCT Pro	AGC Ser	GAT Asp 55	CTT Leu	GTA Val	GGC Gly	ATA Ile	CCG Pro 60	ACT Thr	AGG Arg	ATT Ile	TCC Ser	192
AGG Arg 65	GCA Ala	GAA Glu	GAA Glu	AAG Lys	ATT Ile 70	ACA Thr	TCT Ser	ACA Thr	CTT Leu	GGT Gly 75	TCC Ser	AAT Asn	CAA Gln	GAT Asp	GTA Val 80	240

GTA Val	GAT Asp	AGG Arg	ATA Ile	TAT Tyr 85	AAG Lys	CAA Gln	GTG Val	GCC Ala	CTT Leu 90	GAG Glu	TCT Ser	CCA Pro	TTG Leu	GCA Ala 95	TTG Leu	28	8
TTA Leu	AAT Asn	ACT Thr	GAG Glu 100	ACC Thr	ACA Thr	ATT Ile	ATG Met	AAC Asn 105	GCA Ala	ATA Ile	ACA Thr	TCT Ser	CTC Leu 110	TCT Ser	TAT Tyr	33	6
CAG Gln	ATT Ile	AAT Asn 115	GGA Gly	GCT Ala	GCA Ala	AAC Asn	AAC Asn 120	AGC Ser	GGG Gly	TGG Trp	GGG Gly	GCA Ala 125	CCT Pro	ATT Ile	CAT His	38	4
GAC Asp	CCA Pro 130	GAT Asp	TAT Tyr	ATA Ile	GGG Gly	GGG Gly 135	ATA Ile	GGC Gly	AAA Lys	GAA Glu	CTC Leu 140	ATT Ile	GTA Val	GAT Asp	GAT Asp	43	2
GCT Ala 145	AGT Ser	GAT Asp	GTC Val	ACA Thr	TCA Ser 150	TTC Phe	TAT Tyr	CCC Pro	TCT Ser	GCA Ala 155	TTT Phe	CAA Gln	GAA Glu	CAT His	CTG Leu 160	48	0
AAT Asn	TTT Phe	ATC Ile	CCG Pro	GCG Ala 165	Pro	ACT Thr	ACA Thr	GGA Gly	TCA Ser 170	Gly	TGC Cys	ACT Thr	CGA Arg	ATA Ile 175	Pro	52	8
TCA Ser	TTT Phe	GAC Asp	ATG Met 180	AGT Ser	GCT Ala	ACC Thr	CAT His	TAC Tyr 185	TGC Cys	TAC Tyr	ACC Thr	CAT His	AAT Asn 190	GTA Val	ATA Ile	57	6
			TGC Cys													62	4
			CGG Arg													67	2
			AAC Asn													72	0
			CCC Pro													76	8
ACA Thr	GAG Glu	GAA Glu	GAA Glu 260	GAT Asp	TAT Tyr	AAC Asn	TCA Ser	GCT Ala 265	GTC Val	CCT Pro	ACG Thr	CGG Arg	ATG Met 270	GTA Val	CAT His	81	6
G <b>GG</b> Gly	AGG Arg	TTA Leu 275	GGG Gly	TTC Phe	GAC Asp	GGC Gly	CAA Gln 280	TAT Tyr	CAC His	GAA Glu	AAG Lys	GAC Asp 285	CTA Leu	GAT Asp	GTC Val	86	4
ACA Thr	ACA Thr 290	TTA Leu	TTC Phe	GGG Gly	GAC Asp	TGG Trp 295	GTG Val	GCC Ala	AAC Asn	TAC Tyr	CCA Pro 300	GGA Gly	GTA Val	GGG Gly	GGT Gly	91	2
GGA Gly 305	TCT Ser	TTT Phe	ATT Ile	GAC Asp	AGC Ser 310	CGC Arg	GTG Val	TGG Trp	TTC Phe	TCA Ser 315	GTC Val	TAC Tyr	GGA Gly	GGG Gly	TTA Leu 320	96	0
AAA Lys	CCC Pro	AAT Asn	ACA Thr	CCC Pro 325	AGT Ser	GAC Asp	ACT Thr	GTA Val	CAG Gln 330	GAA Glu	GGG Gly	AAA Lys	TAT Tyr	GTG Val 335	ATA Ile	100	8
			TAC Tyr 340													105	6

CGA Arg	ATG Met	GCC Ala 355	AAG Lys	TCT Ser	TCG Ser	TAT Tyr	AAG Lys 360	CCT Pro	GGA Gly	CGG Arg	TTT Phe	GGT Gly 365	GGG Gly	AAA Lys	CGC Arg	1104
ATA Ile	CAG Gln 370	CAG Gln	GCT Ala	ATC Ile	TTA Leu	TCT Ser 375	ATC Ile	AAA Lys	GTG Val	TCA Ser	ACA Thr 380	TCC Ser	TTA Leu	GGC Gly	GAA Glu	1152
GAC Asp 385	CCG Pro	GTA Val	CTG Leu	ACT Thr	GTA Val 390	CCG Pro	CCC Pro	AAC Asn	ACA Thr	GTC Val 395	ACA Thr	CTC Leu	ATG Met	GGG Gly	GCC Ala 400	1200
GAA Glu	GGC Gly	AGA Arg	ATT Ile	CTC Leu 405	ACA Thr	GTA Val	GGG Gly	ACA Thr	TCC Ser 410	CAT His	TTC Phe	T <b>T</b> G Leu	TAT Tyr	CAG Gln 415	CGA Arg	1248
	TCA Ser															1296
	AAA Lys															1344
	CCA Pro 450															1392
TGT Cys 465	GTT Val	ACT Thr	GGA Gly	GTC Val	TAT Tyr 470	ACA Thr	GAT Asp	CCA Pro	TAT Tyr	CCC Pro 475	CTA Leu	ATC Ile	TTC Phe	TAT Tyr	AGA Arg 480	1440
AAC Asn	CAC His	ACC Thr	TTG Leu	CGA Arg 485	GGG Gly	GTA Val	TTC Phe	GGG Gly	ACA Thr 490	ATG Met	CTT Leu	GAT Asp	GGT Gly	GAA Glu 495	CAA Gln	1488
GCA Ala	AGA Arg	CTT Leu	AAC Asn 500	CCT Pro	GCG Ala	TCT Ser	GCA Ala	GTA Val 505	TTC Phe	GAT Asp	AGC Ser	ACA Thr	TCC Ser 510	CGC Arg	AGT Ser	1536
CGC Arg	ATA Ile	ACT Thr 515	CGA Arg	GTG Val	AGT Ser	TCA Ser	AGC Ser 520	AGC Ser	ATC Ile	AAA Lys	GCA Ala	GCA Ala 525	TAC Tyr	ACA Thr	ACA Thr	1584
TCA Ser	ACT Thr 530	TGT Cys	TTT Phe	AAA Lys	GTG Val	GTC Val 535	AAG Lys	ACC Thr	AAT Asn	AAG Lys	ACC Thr 540	TAT Tyr	TGT Cys	CTC Leu	AGC Ser	1632
ATT Ile 545	GCT Ala	GAA Glu	ATA Ile	TCT Ser	AAT Asn 550	ACT Thr	CTC Leu	TTC Phe	G <b>GA</b> Gly	GAA Glu 555	TTC Phe	AGA Arg	ATC Ile	GTC Val	CCG Pro 560	1680
TTA Leu	CTA Leu	GTT Val	GAG Glu	ATC Ile 565	CTC Leu	AAA Lys	GAT Asp	GAC Asp	GGG Gly 570	GTT Val	AGA Arg	GAA Glu	GCC Ala	AGG Arg 575	TCT Ser	1728
GGC Gly	TAG															1734

# (2) INFORMATION FOR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 577 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear

175

(ii) MOLECULE TYPE: protein

325

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Met Asp Arg Ala Val Ser Gln Val Ala Leu Glu Asn Asp Glu Arg Glu Ala Lys Asn Thr Trp Arg Leu Ile Phe Arg Ile Ala Ile Leu Phe Leu Thr Val Val Thr Leu Ala Ile Ser Val Ala Ser Leu Leu Tyr Ser Met Gly Ala Ser Thr Pro Ser Asp Leu Val Gly Ile Pro Thr Arg Ile Ser Arg Ala Glu Glu Lys Ile Thr Ser Thr Leu Gly Ser Asn Gln Asp Val Val Asp Arg Ile Tyr Lys Gln Val Ala Leu Glu Ser Pro Leu Ala Leu Leu Asn Thr Glu Thr Thr Ile Met Asn Ala Ile Thr Ser Leu Ser Tyr Gln Ile Asn Gly Ala Ala Asn Asn Ser Gly Trp Gly Ala Pro Ile His Asp Pro Asp Tyr Ile Gly Gly Ile Gly Lys Glu Leu Ile Val Asp Asp Ala Ser Asp Val Thr Ser Phe Tyr Pro Ser Ala Phe Gln Glu His Leu Asn Phe Ile Pro Ala Pro Thr Thr Gly Ser Gly Cys Thr Arg Ile Pro Ser Phe Asp Met Ser Ala Thr His Tyr Cys Tyr Thr His Asn Val Ile Leu Ser Gly Cys Arg Asp His Ser His Ser His Gln Tyr Leu Ala Leu Gly Val Leu Arg Thr Ser Ala Thr Gly Arg Val Phe Phe Ser Thr Leu Arg Ser Ile Asn Leu Asp Asp Thr Gln Asn Arg Lys Ser Cys Ser Val 235 Ser Ala Thr Pro Leu Gly Cys Asp Met Leu Cys Ser Lys Ala Thr Glu Thr Glu Glu Glu Asp Tyr Asn Ser Ala Val Pro Thr Arg Met Val His Gly Arg Leu Gly Phe Asp Gly Gln Tyr His Glu Lys Asp Leu Asp Val Thr Thr Leu Phe Gly Asp Trp Val Ala Asn Tyr Pro Gly Val Gly Gly Gly Ser Phe Ile Asp Ser Arg Val Trp Phe Ser Val Tyr Gly Gly Leu Lys Pro Asn Thr Pro Ser Asp Thr Val Glu Glu Lys Tyr Val Ile

Tyr Lys Arg Tyr Asn Asp Thr Cys Pro Asp Glu Gln Asp Tyr Gln Ile 340 345 350

Arg Met Ala Lys Ser Ser Tyr Lys Pro Gly Arg Phe Gly Gly Lys Arg 355 360 365

Ile Gln Gln Ala Ile Leu Ser Ile Lys Val Ser Thr Ser Leu Gly Glu 370 375 380

Asp Pro Val Leu Thr Val Pro Pro Asn Thr Val Thr Leu Met Gly Ala 385 390 395 400

Glu Gly Arg Ile Leu Thr Val Gly Thr Ser His Phe Leu Tyr Gln Arg 405 410 415

Gly Ser Ser Tyr Phe Ser Pro Ala Leu Leu Tyr Pro Met Thr Val Ser 420 425 430

Asn Lys Thr Ala Thr Leu His Ser Pro Tyr Thr Phe Asn Ala Phe Thr 435 440 445

Arg Pro Gly Ser Ile Pro Cys Gln Ala Ser Ala Arg Cys Pro Asn Ser 450 455 460

Cys Val Thr Gly Val Tyr Thr Asp Pro Tyr Pro Leu Ile Phe Tyr Arg 465 470 475 480

Asn His Thr Leu Arg Gly Val Phe Gly Thr Met Leu Asp Gly Glu Gln
485
490
495

Ala Arg Leu Asn Pro Ala Ser Ala Val Phe Asp Ser Thr Ser Arg Ser 500 505 510

Arg Ile Thr Arg Val Ser Ser Ser Ser Ile Lys Ala Ala Tyr Thr Thr 515 520 525

Ser Thr Cys Phe Lys Val Val Lys Thr Asn Lys Thr Tyr Cys Leu Ser 530 535 540

Ile Ala Glu Ile Ser Asn Thr Leu Phe Gly Glu Phe Arg Ile Val Pro 545 550 555 560

Leu Leu Val Glu Ile Leu Lys Asp Asp Gly Val Arg Glu Ala Arg Ser 565 570 575

Gly

#### (2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1662 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..1662
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

ATG Met 1	GGC Gly	TCC Ser	AGA Arg	CCT Pro 5	TCT Ser	ACC Thr	AAG Lys	AAC Asn	CCA Pro 10	GCA Ala	CCT Pro	ATG Met	ATG Met	CTG Leu 15	ACT Thr		48
										TGT Cys						9	96
GAT Asp	GGC Gly	AGG Arg 35	CCT Pro	CTT Leu	GCA Ala	GCT Ala	GCA Ala 40	GGA Gly	ATT Ile	GTG Val	GTT Val	ACA Thr 45	GGA Gly	GAC Asp	AAA Lys	1	44
GCA Ala	GTC Val 50	AAC Asn	ATA Ile	TAC Tyr	ACC Thr	TCA Ser 55	TCC Ser	CAG Gln	ACA Thr	GGA Gly	TCA Ser 60	ATC Ile	ATA Ile	GTT Val	AAG Lys	1	92
CTC Leu 65	CTC Leu	CCG Pro	AAT Asn	CTG Leu	CCA Pro 70	AAG Lys	GAT Asp	AAG Lys	GAG Glu	GCA Ala 75	TGT Cys	GCG Ala	AAA Lys	GCC Ala	CCC Pro 80	2	240
TTG Leu	GAT Asp	GCA Ala	TAC Tyr	AAC Asn 85	AGG Arg	ACA Thr	TTG Leu	ACC Thr	ACT Thr 90	TTG Leu	CTC Leu	ACC Thr	CCC Pro	CTT Leu 95	GGT Gly	2	288
GAC Asp	TCT Ser	ATC Ile	CGT Arg 100	AGG Arg	ATA Ile	CAA Gln	GAG Glu	TCT Ser 105	GTG Val	ACT Thr	ACA Thr	TCT Ser	GGA Gly 110	GGG Gly	GGG Gly	3	336
AGA Arg	CAG Gln	GGG Gly 115	CGC Arg	CTT Leu	ATA Ile	GGC Gly	GCC Ala 120	ATT Ile	ATT Ile	GGC Gly	GGT Gly	GTG Val 125	GCT Ala	CTT Leu	GGG Gly	3	384
GTT Val	GCA Ala 130	ACT Thr	GCC Ala	GCA Ala	CAA Gln	ATA Ile 135	ACA Thr	GCG Ala	GCC Ala	GCA Ala	GCT Ala 140	CTG Leu	ATA Ile	CAA Gln	GCC Ala	4	432
AAA Lys 145	CAA Gln	AAT Asn	GCT Ala	GCC Ala	AAC Asn 150	ATC Ile	CTC Leu	CGA Arg	CTT Leu	AAA Lys 155	GAG Glu	AGC Ser	ATT Ile	GCC Ala	GCA Ala 160	•	480
Thr	Asn	Glu	Ala	Val 165	His	Glu	Val	Thr	170	GIĀ	ьeu	ser	GIII	175	GCA Ala	!	528
GTG Val	GCA Ala	GTT Val	GGG Gly 180	AAG Lys	ATG Met	CAG Gln	CAG Gln	TTC Phe 185	GTT Val	AAT Asn	GAC Asp	CAA Gln	TTT Phe 190	AAT Asn	AAA Lys	!	<b>57</b> 6
ACA Thr	GCT Ala	CAG Gln 195	Glu	TTA Leu	GAC Asp	TGC Cys	ATC Ile 200	Lys	ATT	GCA Ala	CAG Gln	CAA Gln 205	GTT Val	GGT Gly	GTA Val	,	624
GAG Glu	CTC Leu 210	Asn	CTG Leu	TAC Tyr	CTA Leu	ACC Thr 215	GAA Glu	TCG Ser	ACT Thr	ACA Thr	GTA Val 220	Pne	GGA Gly	CCA Pro	CAA Gln		672
ATC Ile 225	ACT Thr	TCA Ser	CCT Pro	GCC Ala	TTA Leu 230	Asn	AAG Lys	CTG Leu	ACT Thr	ATT Ile 235	Gin	GCA Ala	CTT Leu	TAC	AAT Asn 240		720
CTA Leu	GCT Ala	GGT Gly	GGG Gly	AAT Asn 245	Met	GAT Asp	TAC	TTA Leu	TTG Leu 250	Thr	AAG Lys	TTA Leu	GGT Gly	ATA Ile 255	GGG Gly		768
<b>AA</b> C Asn	AAT Asn	CAA Gln	CTC Leu 260	Ser	TCA Ser	TTA Leu	ATC Ile	GGT Gly 265	Ser	GGC Gly	TTA Leu	ATC Ile	ACC Thr 270	GIY	AAC Asn		816

									CTC Leu							864
									ATG Met							912
									TTT Phe							960
									ATA Ile 330							1008
									TAT Tyr							1056
									TGC <b>Cy</b> s							1104
									GCA Ala							1152
									TGC Cys							1200
									CAA Gln 410							1248
TCT Ser	CTA Leu	ATA Ile	GAT Asp 42	Lys	CAA Gln	TCA Ser	TGC Cys	AAT Asn 429	GTT Val	TTA Leu	TCC Ser	TTA Leu	GGC Gly 430	Gly	ATA Ile	1296
ACT Thr	TTA Leu	AGG Arg 435	CTC Leu	AGT Ser	GGG Gly	GAA Glu	TTC Phe 440	GAT Asp	GTA Val	ACT Thr	TAT Tyr	CAG Gln 445	AAG Lys	AAT Asn	ATC Ile	1344
TCA Ser	ATA Ile 450	CAA Gln	GAT Asp	TCT Ser	CAA Gln	GTA Val 455	ATA Ile	ATA Ile	ACA Thr	GGC Gly	AAT Asn 460	CTT Leu	GAT Asp	ATC Ile	TCA Ser	1392
ACT Thr 465	GAG Glu	CTT Leu	GGG Gly	AAT Asn	GTC Val 470	AAC Asn	AAC Asn	TCG Ser	ATC Ile	AGT Ser 475	AAT Asn	GCC Ala	TTG Leu	AAT Asn	AAG Lys 480	1440
TTA Leu	GAG Glu	GAA Glu	AGC Ser	AAC Asn 485	AGA Arg	AAA Lys	CTA Leu	GAC Asp	AAA Lys 490	GTC Val	AAT Asn	GTC Val	AAA Lys	CTG Leu 495	ACC Thr	1488
AGC Ser	ACA Thr	TCT Ser	GCT Ala 500	CTC Leu	ATT Ile	ACC Thr	TAT Tyr	ATC Ile 505	GTT Val	TTG Leu	ACT Thr	ATC Ile	ATA Ile 510	TCT Ser	CTT Leu	1536
GT <b>T</b> Val	TTT Phe	GGT Gly 515	ATA Ile	CTT Leu	AGC Ser	CTG Leu	ATT Ile 520	CTA Leu	GCA Ala	TGC Cys	TAC Tyr	CTA Leu 525	ATG Met	TAC Tyr	AAG Lys	1584
CAA Gln	AAG Lys 530	GCG Ala	CAA Gln	CAA Gln	AAG Lys	ACC Thr 535	TTA Leu	TTA Leu	TGG Trp	CTT Leu	GGG Gly 540	AAT Asn	AAT Asn	ACC Thr	CTA Leu	1632

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179

GAT CAG ATG AGA GCC ACT ACA AAA ATG TGA Asp Gln Met Arg Ala Thr Thr Lys Met 550

1662

## (2) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 553 amino acids (B) TYPE: amino acid

  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13: Met Gly Ser Arg Pro Ser Thr Lys Asn Pro Ala Pro Met Met Leu Thr Ile Arg Val Ala Leu Val Leu Ser Cys Ile Cys Pro Ala Asn Ser Ile Asp Gly Arg Pro Leu Ala Ala Gly Ile Val Val Thr Gly Asp Lys Ala Val Asn Ile Tyr Thr Ser Ser Gln Thr Gly Ser Ile Ile Val Lys Leu Leu Pro Asn Leu Pro Lys Asp Lys Glu Ala Cys Ala Lys Ala Pro Leu Asp Ala Tyr Asn Arg Thr Leu Thr Thr Leu Leu Thr Pro Leu Gly Asp Ser Ile Arg Arg Ile Gln Glu Ser Val Thr Thr Ser Gly Gly 105 Arg Gln Gly Arg Leu Ile Gly Ala Ile Ile Gly Gly Val Ala Leu Gly Val Ala Thr Ala Ala Gln Ile Thr Ala Ala Ala Ala Leu Ile Gln Ala 135 Lys Gln Asn Ala Ala Asn Ile Leu Arg Leu Lys Glu Ser Ile Ala Ala 145 Thr Asn Glu Ala Val His Glu Val Thr Asp Gly Leu Ser Gln Leu Ala Val Ala Val Gly Lys Met Gln Gln Phe Val Asn Asp Gln Phe Asn Lys 185 Thr Ala Gln Glu Leu Asp Cys Ile Lys Ile Ala Gln Gln Val Gly Val Glu Leu Asn Leu Tyr Leu Thr Glu Ser Thr Thr Val Phe Gly Pro Gln 220 215 Ile Thr Ser Pro Ala Leu Asn Lys Leu Thr Ile Gln Ala Leu Tyr Asn

Leu Ala Gly Gly Asn Met Asp Tyr Leu Leu Thr Lys Leu Gly Ile Gly

Asn Asn Gln Leu Ser Ser Leu Ile Gly Ser Gly Leu Ile Thr Gly Asn 265

245

Pro Ile Leu Tyr Asp Ser Gln Thr Gln Leu Leu Gly Ile Gln Val Thr 280

Leu Pro Ser Val Gly Asn Leu Asn Asn Met Arg Ala Thr Tyr Leu Glu 290

Thr Leu Ser Val Ser Thr Thr Arg Gly Phe Ala Ser Ala Leu Val Pro 305 310 315 320

Lys Val Val Thr Arg Val Gly Ser Val Ile Glu Glu Leu Asp Thr Ser 325 330 335

Tyr Cys Ile Glu Thr Asp Leu Asp Leu Tyr Cys Thr Arg Ile Val Thr 340 345 350

Phe Pro Met Ser Pro Gly Ile Tyr Ser Cys Leu Ser Gly Asn Thr Ser 355 360 365

Ala Cys Met Tyr Ser Lys Thr Glu Gly Ala Leu Thr Thr Pro Tyr Met 370 375 380

Thr Ile Lys Gly Ser Val Ile Ala Asn Cys Lys Met Thr Thr Cys Arg 385 390 395

Cys Val Asn Pro Pro Gly Ile Ile Ser Gln Asn Tyr Gly Glu Ala Val 405 410 415

Ser Leu Ile Asp Lys Gln Ser Cys Asn Val Leu Ser Leu Gly Gly Ile 420 425 430

Thr Leu Arg Leu Ser Gly Glu Phe Asp Val Thr Tyr Gln Lys Asn Ile 435 440 445

Ser Ile Gln Asp Ser Gln Val Ile Ile Thr Gly Asn Leu Asp Ile Ser 450 455 460

Thr Glu Leu Gly Asn Val Asn Asn Ser Ile Ser Asn Ala Leu Asn Lys
465 470 475 480

Leu Glu Glu Ser Asn Arg Lys Leu Asp Lys Val Asn Val Lys Leu Thr 485 490 495

Ser Thr Ser Ala Leu Ile Thr Tyr Ile Val Leu Thr Ile Ile Ser Leu 500 505 510

Val Phe Gly Ile Leu Ser Leu Ile Leu Ala Cys Tyr Leu Met Tyr Lys 515 520 525

Gln Lys Ala Gln Gln Lys Thr Leu Leu Trp Leu Gly Asn Asn Thr Leu 530 540

Asp Gln Met Arg Ala Thr Thr Lys Met 545

# (2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 3489 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO

181

(iv) ANTI-SENSE: NO

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1..3489

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

	(xi	) SE(	QUEN	CE DI	ESCR:	IPTIC	ON: S	SEQ I	ID NO	0:14:	:						
														CTA Leu 15			48
														CAA Gln		!	96
														TAT Tyr		14	44
		AAT					TCT							CCT Pro		1	92
														TCT Ser		24	40
														AGT Ser 95		28	88
														GTT Val		3:	36
CAT His	TGT Cys	TAT Tyr 115	AAA Lys	TAT Tyr	GAT Asp	GGG Gly	TGT Cys 120	CCT Pro	ATA Ile	ACT Thr	GGC Gly	ATG Met 125	CTT Leu	CAA Gln	AAG Lys	31	84
AAT Asn	TTT Phe 130	Leu	CGT Arg	GTT Val	TCT Ser	GCT Ala 13	Met	AAA Lys	AAT Asn	GGC Gly	CAG Gln 140	Leu	TTC Phe	TAT Tyr	AAT Asn	4:	32
TTA Leu 145	ACA Thr	GTT Val	AGT Ser	GTA Val	GCT Ala 150	AAG Lys	TAC Tyr	CCT Pro	ACT Thr	TTT Phe 155	AAA Lys	TCA Ser	TTT Phe	CAG Gln	TGT Cys 160	4:	80
GTT Val	AAT Asn	AAT Asn	TTA Leu	ACA Thr 165	TCC Ser	GTA Val	TAT Tyr	TTA Leu	AAT Asn 170	GGT Gly	GAT Asp	CTT Leu	GTT Val	TAC Tyr 175	ACC Thr	5:	28
TCT Ser	AAT Asn	GAG Glu	ACC Thr 180	ACA Thr	GAT Asp	GTT Val	ACA Thr	TCT Ser 185	GCA Ala	GGT Gly	GTT Val	TAT Tyr	TTT Phe 190	AAA Lys	GCT Ala	5	76
GGT Gly	GGA Gly	CCT Pro 195	ATA Ile	ACT Thr	TAT Tyr	AAA Lys	GTT Val 200	ATG Met	AGA Arg	AAA Lys	GTT Val	AAA Lys 205	GCC Ala	CTG Leu	GCT Ala	6:	24
TAT Tyr	TTT Phe 210	GTT Val	AAT Asn	GGT Gly	ACT Thr	GCA Ala 215	CAA Gln	GAT Asp	GTT Val	ATT Ile	TTG Leu 220	TGT Cys	GAT Asp	GGA Gly	TCA Ser	6	72
CCT Pro 225	AGA Arg	GGC Gly	TTG Leu	TTA Leu	GCA Ala 230	TGC Cys	CAG Gln	TAT Tyr	AAT Asn	ACT Thr 235	GGC Gly	AAT Asn	TTT Phe	TCA Ser	GAT Asp 240	7.	20

GGC Gly	TTT Phe	TAT	CCT Pro	TTT Phe 245	Ile	AAT Asn	AGT Ser	AGT Ser	TTA Leu 250	GTT Val	AAG Lys	CAG Gln	AAG Lys	TTT Phe 255	ATT Ile	768
GTC Val	TAT Tyr	CGT Arg	GAA Glu 260	AAT Asn	AGT Ser	GTT Val	AAT Asn	ACT Thr 265	ACT Thr	TTT Phe	ACG Thr	TTA Leu	CAC His 270	AAT Asn	TTC Phe	816
ACT Thr	TTT Phe	CAT His 275	AAT Asn	GAG Glu	ACT Thr	GGC Gly	GCC Ala 280	AAC Asn	CCT Pro	AAT Asn	CCT Pro	AGT Ser 285	GGT Gly	GTT Val	CAG Gln	864
AAT Asn	ATT Ile 290	Leu	ACT Thr	TAC Tyr	CAA Gln	ACA Thr 295	CAA Gln	ACA Thr	GCT Ala	CAG Gln	AGT Ser 300	GGT Gly	TAT Tyr	TAT Tyr	AAT Asn	912
			TCC Ser													960
ATG Met	TAT Tyr	GGA Gly	TCT Ser	TAT Tyr 325	CAC His	CCA Pro	AGT Ser	TGT Cys	AAT Asn 330	TTT Phe	AGA Arg	CTA Leu	GAA Glu	ACT Thr 335	ATT Ile	1008
			TTG Leu 340													1056
CCT Pro	CTT Leu	CAA Gln 355	GGT Gly	GGT Gly	TGC Cys	AAG Lys	CAA Gln 360	TCT Ser	GTC Val	TTT Phe	AGT Ser	GGT Gly 365	AGA Arg	GCA Ala	ACT Thr	1104
TGT Cys	TGT Cys 370	TAT Tyr	GCT Ala	TAT Tyr	TCA Ser	TAT Tyr 375	GGA Gly	GGT Gly	CCT Pro	TCG Ser	CTG Leu 380	TGT Cys	AAA Lys	GGT Gly	GTT Val	1152
TAT Tyr 385	TCA Ser	GGT Gly	GAG Glu	TTA Leu	GAT Asp 390	CTT Leu	AAT Asn	TTT Phe	GAA Glu	TGT Cys 395	GGA Gly	CTG Leu	TTA Leu	GTT Val	TAT Tyr 400	1200
GTT Val	ACT Thr	AAG Lys	AGC Ser	GGT Gly 405	GGC Gly	TCT Ser	CGT Arg	ATA Ile	CAA Gln 410	ACA Thr	GCC Ala	ACT Thr	GAA Glu	CCG Pro 415	CCA Pro	1248
GTT Val	ATA Ile	ACT Thr	CGA Arg 420	CAC His	AAT Asn	TAT Tyr	AAT Asn	AAT Asn 425	ATT Ile	ACT Thr	TTA Leu	AAT Asn	ACT Thr 430	TGT Cys	GTT Val	1296
GAT Asp	TAT Tyr	AAT Asn 435	ATA Ile	TAT Tyr	GGC Gly	AGA Arg	ACT Thr 440	GGC Gly	CAA Gln	GGT Gly	TTT Phe	ATT Ile 445	ACT Thr	AAT Asn	GTA Val	1344
ACC Thr	GAC Asp 450	TCA Ser	GCT Ala	GTT Val	AGT Ser	TAT Tyr 455	AAT Asn	TAT Tyr	CTA Leu	GCA Ala	GAC Asp 460	GCA Ala	GGT Gly	TTG Leu	GCT Ala	1392
ATT Ile 465	TTA Leu	GAT Asp	ACA Thr	TCT Ser	GGT Gly 470	TCC Ser	ATA Ile	GAC Asp	ATC Ile	TTT Phe 475	GTT Val	GTA Val	CAA Gln	GGT Gly	GAA Glu 480	1440
TAT Tyr	GGT Gly	CTT Leu	ACT Thr	TAT Tyr 485	TAT Tyr	AAG Lys	GTT Val	AAC Asn	CCT Pro 490	TGC Cys	GAA Glu	GAT Asp	GTC Val	AAC Asn 495	CAG Gln	1488
CAG Gln	TTT Phe	GTA Val	GTT Val 500	TCT Ser	GGT Gly	GGT Gly	AAA Lys	TTA Leu 505	GTA Val	GGT Gly	ATT Ile	CTT Leu	ACT Thr 510	TCA Ser	CGT Arg	1536

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							CTT Leu 520									1584
							AGA Arg									1632
							GGT Gly									1680
							AAA Lys									1728
							GTG Val									1776
							CAA Gln 600									1824
AAT Asn	TGT Cys 610	CTG Leu	CAG Gln	TAT Tyr	GTT Val	TGT Cys 615	GGC Gly	AAT Asn	TCT Ser	CTG Leu	GAT Asp 620	TGT Cys	AGA Arg	GAT Asp	TTG Leu	1872
TTT Phe 625	CAA Gln	CAA Gln	TAT Tyr	GGG Gly	CCT Pro 630	GTT Val	TGT Cys	GAC Asp	AAC Asn	ATA Ile 635	TTG Leu	TCT Ser	GTA Val	GTA Val	AAT Asn 640	1920
AGT Ser	ATT Ile	GGT Gly	CAA Gln	AAA Lys 645	GAA Glu	GAT Asp	ATG Met	GAA Glu	CTT Leu 650	TTG Leu	AAT Asn	TTC Phe	TAT Tyr	TCT Ser 655	TCT Ser	1968
ACT Thr	AAA Lys	CCG Pro	GCT Ala 660	GGT Gly	TTT Phe	AAT Asn	ACA Thr	CCA Pro 665	TTT Phe	CTT Leu	AGT Ser	AAT Asn	GTT Val 670	AGC Ser	ACT Thr	2016
GGT Gly	GAG Glu	TTT Phe 675	AAT Asn	ATT Ile	TCT Ser	CTT Leu	CTG Leu 680	TTA Leu	ACA Thr	ACT Thr	CCT Pro	AGT Ser 685	AGT Ser	CCT Pro	AGA Arg	2064
AGG Arg	CGT Arg 690	TCT Ser	TTT Phe	ATT Ile	GAA Glu	GAC Asp 695	CTT Leu	CTA Leu	TTT Phe	ACA Thr	AGC Ser 700	GTT Val	GAA Glu	TCT Ser	GTT Val	2112
GGA Gly 705	TTA Leu	CCA Pro	ACA Thr	GAT Asp	GAC Asp 710	GCA Ala	TAC Tyr	AAA Lys	AAT Asn	TGC Cys 715	ACT Thr	GCA Ala	GGA Gly	CCT Pro	TTA Leu 720	2160
GGT Gly	TTT Phe	CTT Leu	AAG Lys	GAC Asp 72	Leu	GCG Ala	TGT Cys	GCT Ala	CGT Arg 73	Glu	TAT Tyr	AAT Asn	GGT Gly	TTG Leu 73	Leu	2208
GTG Val	TTG Leu	CCT Pro	CCC Pro 740	ATT Ile	ATA Ile	ACA Thr	GCA Ala	GAA Glu 745	ATG Met	CAA Gln	ACT Thr	TTG Leu	TAT Tyr 750	ACT Thr	AGT Ser	2256
TCT Ser	CTA Leu	GTA Val 755	GCT Ala	TCT Ser	ATG Met	GCT Ala	TTT Phe 760	GGT Gly	GGT Gly	ATT Ile	ACT Thr	GCA Ala 765	GCT Ala	GGT Gly	GCT Ala	2304
ATA Ile	CCT Pro 770	TT <b>T</b> Phe	GCC Ala	ACA Thr	CAA Gln	CTG Leu 775	CAG Gln	GCT Ala	AGA Arg	ATT Ile	AAT Asn 780	CAC His	TTG Leu	GGT Gly	ATT Ile	2352

					TTG Leu 790											2400
					CGT Arg											2448
GCA Ala	TTA Leu	CAA Gln	CAA Gln 820	ATT Ile	CAA Gln	GAT Asp	GTT Val	GTT Val 825	AAT Asn	AAG Lys	CAG Gln	AGT Ser	GCT Ala 830	ATT Ile	CTT Leu	2496
ACT Thr	GAG Glu	ACT Thr 835	ATG Met	GCA Ala	TCA Ser	CTT Leu	AAT Asn 840	AAA Lys	AAT Asn	TTT Phe	GGT Gly	GCT Ala 845	ATT Ile	TCT Ser	TCT Ser	2544
GTG Val	ATT Ile 850	CAA Gln	GAA Glu	ATC Ile	TAC Tyr	CAG Gln 855	CAA Gln	CTT Leu	GAC Asp	GCC Ala	ATA Ile 860	CAA Gln	GCA Ala	AAT Asn	GCT Ala	2592
CAA Gln 865	GTG Val	GAT Asp	CGT Arg	CTT Leu	ATA Ile 870	ACT Thr	GGT Gly	AGA Arg	TTG Leu	TCA Ser 875	TCA Ser	CTT Leu	TCT Ser	GTT Val	TTA Leu 880	2640
GCA Ala	TCT Ser	GCT Ala	AAG Lys	CAG Gln 885	GCG Ala	GAG Glu	CAT His	ATT Ile	AGA Arg 890	GTG Val	TCA Ser	CAA Gln	CAG Gln	CGT Arg 895	GAG Glu	2688
TTA Leu	GCT Ala	ACT Thr	CAG Gln 900	AAA Lys	ATT Ile	AAT Asn	GAG Glu	TGT Cys 905	GTT Val	AAG Lys	TCA Ser	CAG Gln	TCT Ser 910	ATT Ile	AGG Arg	2736
TAC Tyr	TCC Ser	TTT Phe 915	TGT Cys	GGT Gly	AAT Asn	GGA Gly	CGA Arg 920	CAT His	GTT Val	CTA Leu	ACC Thr	ATA Ile 925	CCG Pro	CAA Gln	AAT Asn	2784
					GTG Val											2832
					GCA Ala 950											2880
GCT Ala	AGT Ser	CAG Gln	TAT Tyr	GCA Ala 965	ATA Ile	GTA Val	CCC Pro	GCT Ala	AAT Asn 970	GGT Gly	AGG Arg	GGT Gly	ATT Ile	TTT Phe 975	ATA Ile	2928
					TAC Tyr											2976
					GGA Gly			Val					Cys			3024
		Val			AAT Asn		Thr					Phe				3072
	Asp				AAT Asn 1030	Asp					Trp					3120
					GAC Asp					Asn					Ile	3168

									185								
									103								
CTT Leu	GAC Asp	ATT Ile	GAT Asp 106	Ser	GAA Glu	ATT Ile	GAT Asp	CGT Arg 106	Ile	CAA Gln	GGC Gly	GTT Val	ATA Ile 107	Gln	GGT Gly	3	3216
CTT Leu	AAT Asn	GAC Asp 10	Ser	TTA Leu	ATA Ile	GAC Asp	CTT Leu 108	Glu	AAA Lys	CTT Leu	TCA Ser	ATA Ile 108	Leu	AAA Lys	ACT Thr	3	3264
		AAG Lys 0					Val	_				Ala				3	3312
	Ile	TTC Phe				Leu					Phe					3	360
		TGT Cys			Gly					Met					Lys	3	1408
		AAG Lys		Ser					Thr					Val		3	3456
		CAA Gln 1155	Asn					Ser		TAA						3	1489
(2)	INF	ORMAT	rion	FOR	SEQ	ID N	NO:15	5:									
		(i) S	(A) (B)	LEI	NGTH:		52 ar	nino id		is							
	(:	ii) N	OLE	TULE	TYPI	E: pi	rote:	in									
	()	xi) a	EQUI	ENCE	DESC	CRIPT	NOIT	: SE(	Q ID	NO: 1	L5:						
	_				_	_				T		C	37-7	T	Caro		

Met leu Val Thr Pro Leu Leu Leu Val Thr Leu Cys Val Leu Cys
Ser Ala Ala Leu Tyr Asp Ser Ser Ser Tyr Val Tyr Tyr Gln Ser
Ala Phe Arg 35 Pro Pro Asn Gly Trp His Leu His Gly Gly Ala Tyr Ala
Val Val Asn Ile Ser Ser Glu Ser Asn Asn Ala Gly Ser Ser Pro Gly
65 Ile Val Gly Thr Ile His Gly Gly Arg Val Val Asn Ala Ser Ser
Ala Met Thr Ala Pro Ser Ser Gly Met Ala Trp Ser Ser Ser Gln
Phe Cys Thr Ala His Cys Asn Phe Ser Asp Thr Thr Val Phe Val Thr
His Cys Tyr Lys Tyr Asp Gly Cys Pro Ile Thr Gly Met Leu Gln Lys

Asn Phe Leu Arg Val Ser Ala Met Lys Asn Gly Gln Leu Phe Tyr Asn

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Let 145	Thi	r Val	l Ser	. Val	150	Lys	з Туг	Pro	Thr	155	Lys	s Ser	Phe	Glr	1 Cyr
Val	Asn	n Asn	Leu	165		Val	Туг	Lev	170		/ Asp	Let	ı Val	Ty:	
Ser	Asn	Glu	Thr 180	Thr	Asp	• Val	. Thr	Ser 185		Gly	⁄ Val	Tyr	Phe 190	_	Ala
Gly	Gly	Pro 195		Thr	Tyr	Lys	Val 200	Met	Arg	Lys	: Val	Lys 205		Leu	ı Ala
Tyr	Phe 210	Val	Asn	Gly	Thr	Ala 215		Asp	Val	Ile	220		Asp	Gly	sei
Pro 225	Arg	Gly	Leu	Leu	Ala 230		Gln	Tyr	Asn	Thr 235		' Asn	Phe	Ser	Asp 240
Gly	Phe	Tyr	Pro	Phe 245		Asn	Ser	Ser	Leu 250		Lys	Gln	Lys	Phe 255	
Val	Tyr	Arg	Glu 260		Ser	Val	Asn	Thr 265	Thr	Phe	Thr	Leu	His 270		Phe
Thr	Phe	His 275	Asn	Glu	Thr	Gly	Ala 280	Asn	Pro	Asn	Pro	Ser 285		Val	Glr
Asn	Ile 29	Leu 0	Thr	Tyr	Gln	Thr 29		Thr	Ala	Gln	Ser 30		Tyr	Tyr	Asr
Phe 305	Asn	Phe	Ser	Phe	Leu 310	Ser	Ser	Phe	Val	Tyr 315		Glu	Ser	Asn	Phe 320
Met	Tyr	Gly	Ser	Tyr 325	His	Pro	Ser	Cys	Asn 330	Phe	Arg	Leu	Glu	Thr 335	
Asn	Asn	Gly	Leu 340	Trp	Phe	Asn	Ser	Leu 345	Ser	Val	Ser	Ile	Ala 350	Tyr	Gly
Pro	Leu	Gln 3 <b>5</b> 5	Gly	Gly	Cys	Lys	Gln 360	Ser	Val	Phe	Ser	Gly 365	Arg	Ala	Thr
Cys	Cys 370	Tyr	Ala	Tyr	Ser	Tyr 375	Gly	Gly	Pro	Ser	Leu 380	Cys	Lys	Gly	Val
Tyr 385	Ser	Gly	Glu	Leu	Asp 390	Leu	Asn	Phe	Glu	Cys 395	Gly	Leu	Leu	Val	Tyr 400
Val	Thr	Lys	Ser	Gly 405	Gly	Ser	Arg	Ile	Gln 410	Thr	Ala	Thr	Glu	Pro 415	Pro
Val	Ile	Thr	Arg 420	His	Asn	Tyr	Asn	Asn 425	Ile	Thr	Leu	Asn	Thr 430	Cys	Val
Asp	Tyr	Asn 435	Ile	Tyr	Gly	Arg	Thr 440	Gly	Gln	Gly	Phe	Ile 445	Thr	Asn	Val
Thr	Asp 450	Ser	Ala	Val	Ser	Tyr 455	Asn	Tyr	Leu	Ala	Asp 460	Ala	Gly	Leu	Ala
Ile 465	Leu	Asp	Thr	Ser	Gly 470	Ser	Ile	Asp	Ile	Phe 475	Val	Val	Gln	Gly	Glu 480
Tyr	Gly	Leu	Thr	Tyr 485	Tyr	Lys	Val	Asn	Pro	Cys	Glu	Asp	Val	Asn	Gln

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Gln Phe Val Val Ser Gly Gly Lys Leu Val Gly Ile Leu Thr Ser Arg Asn Glu Thr Gly Ser Gln Leu Leu Glu Asn Gln Phe Tyr Ile Lys Ile 520 Thr Asn Gly Thr Arg Arg Phe Arg Arg Ser Ile Thr Glu Asn Val Ala Asn Cys Pro Tyr Val Ser Tyr Gly Lys Phe Cys Ile Lys Pro Asp Gly Ser Ile Ala Thr Ile Val Pro Lys Gln Leu Glu Gln Phe Val Ala Pro 565 570 Leu Leu Asn Val Thr Glu Asn Val Leu Ile Pro Asn Ser Phe Asn Leu 585 Thr Val Thr Asp Glu Tyr Ile Gln Thr Arg Met Asp Lys Val Gln Ile Asn Cys Leu Gln Tyr Val Cys Gly Asn Ser Leu Asp Cys Arg Asp Leu Phe Gln Gln Tyr Gly Pro Val Cys Asp Asn Ile Leu Ser Val Val Asn Ser Ile Gly Gln Lys Glu Asp Met Glu Leu Leu Asn Phe Tyr Ser Ser Thr Lys Pro Ala Gly Phe Asn Thr Pro Phe Leu Ser Asn Val Ser Thr Gly Glu Phe Asn Ile Ser Leu Leu Leu Thr Thr Pro Ser Ser Pro Arg 680 Arg Arg Ser Phe Ile Glu Asp Leu Leu Phe Thr Ser Val Glu Ser Val Gly Leu Pro Thr Asp Asp Ala Tyr Lys Asn Cys Thr Ala Gly Pro Leu Gly Phe Leu Lys Asp Leu Ala Cys Ala Arg Glu Tyr Asn Gly Leu Leu 730 Val Leu Pro Pro Ile Ile Thr Ala Glu Met Gln Thr Leu Tyr Thr Ser Ser Leu Val Ala Ser Met Ala Phe Gly Gly Ile Thr Ala Ala Gly Ala Ile Pro Phe Ala Thr Gln Leu Gln Ala Arg Ile Asn His Leu Gly Ile Thr Gln Ser Leu Leu Leu Lys Asn Gln Glu Lys Ile Ala Ala Ser Phe Asn Lys Ala Ile Gly Arg Met Gln Glu Gly Phe Arg Ser Thr Ser Leu 810 Ala Leu Gln Gln Ile Gln Asp Val Val Asn Lys Gln Ser Ala Ile Leu 825 Thr Glu Thr Met Ala Ser Leu Asn Lys Asn Phe Gly Ala Ile Ser Ser 840

Val Ile Gln Glu Ile Tyr Gln Gln Leu Asp Ala Ile Gln Ala Asn Ala 850 855 860

Gln Val Asp Arg Leu Ile Thr Gly Arg Leu Ser Ser Leu Ser Val Leu 865 870 875 880

Ala Ser Ala Lys Gln Ala Glu His Ile Arg Val Ser Gln Gln Arg Glu 885 890 895

Leu Ala Thr Gln Lys Ile Asn Glu Cys Val Lys Ser Gln Ser Ile Arg 900 905 910

Tyr Ser Phe Cys Gly Asn Gly Arg His Val Leu Thr Ile Pro Gln Asn 915 920 925

Ala Pro Asn Gly Ile Val Phe Ile His Phe Ser Tyr Thr Pro Asp Ser 930 935 940

Phe Val Asn Val Thr Ala Ile Val Gly Phe Cys Val Lys Pro Ala Asn 945 950 955 960

Ala Ser Gln Tyr Ala Ile Val Pro Ala Asn Gly Arg Gly Ile Phe Ile 965 970 975

Gln Val Asn Gly Ser Tyr Tyr Ile Thr Ala Arg Asp Met Tyr Met Pro 980 985 990

Arg Ala Ile Thr Ala Gly Asp Ile Val Thr Leu Thr Ser Cys Gln Ala 995 1000 1005

Asn Tyr Val Ser Val Asn Lys Thr Val Ile Thr Thr Phe Val Asp Asn 1010 1015 1020

Asp Asp Phe Asp Phe Asn Asp Glu Leu Ser Lys Trp Trp Asn Asp Thr 1025 1030 1035 1040

Lys His Glu Leu Pro Asp Phe Asp Lys Phe Asn Tyr Thr Val Pro Ile 1045 1050 1055

Leu Asp Ile Asp Ser Glu Ile Asp Arg Ile Gln Gly Val Ile Gln Gly
1060 1065 1070

Leu Asn Asp Ser Leu Ile Asp Leu Glu Lys Leu Ser Ile Leu Lys Thr 1075 1080 1085

Tyr Ile Lys Trp Pro Trp Tyr Val Trp Leu Ala Ile Ala Phe Ala Thr 1090 1095 1100

Ile Ile Phe Ile Leu Ile Leu Gly Trp Val Phe Phe Met Thr Gly Cys 1105 1110 1115 1120

Cys Gly Cys Cys Cys Gly Cys Phe Gly Ile Met Pro Leu Met Ser Lys 1125 1130 1135

Cys Gly Lys Lys Ser Ser Tyr Tyr Thr Thr Phe Asp Asn Asp Val Val 1140 1145 1150

Thr Glu Gln Asn Arg Pro Lys Lys Ser Val 1155 1160

## (2) INFORMATION FOR SEQ ID NO:16:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1846 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double

189 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (iii) HYPOTHETICAL: NO (iv) ANTI-SENSE: NO (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..1846 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: ATG TTG GTG AAG TCA CTG TTT CTA GTG ACC ATT TTG TTT GCA CTA TGT 48 Met Leu Val Lys Ser Leu Phe Leu Val Thr Ile Leu Phe Ala Leu Cys AGT GCT AAT TTA TAT GAC AAC GAA TCT TTT 3TG TAT TAC TAC CAG AGT 96 Ser Ala Asn Leu Tyr Asp Asn Glu Ser Phe Cal Tyr Tyr Gln Ser 25 GCT TTT AGG CCA GGA CAT GGT TGG CAT TTA CAT GGA GGT GCT TAT GCA 144 Ala Phe Arg Pro Gly His Gly Trp His Leu His Gly Gly Ala Tyr Ala 40 GTA GTT AAT GTG TCT AGT GAA AAT AAT AAT GCA GGT ACT GCC CCA AGT 192 Val Val Asn Val Ser Ser Glu Asn Asn Ala Gly Thr Ala Pro Ser TGC ACT GCT GCT ATT GGC TAC AGT AAG AAT TTC AGT GCG GCC TCA 240 Cys Thr Ala Gly Ala Ile Gly Tyr Ser Lys Asn Phe Ser Ala Ala Ser 70 GTA GCC ATG ACT GCA CCA CTA AGT GGT ATG TCA TGG TCT GCC TCA TCT Val Ala Met Thr Ala Pro Leu Ser Gly Met Ser Trp Ser Ala Ser Ser 85 TTT TGT ACA GCT CAC TGT AAT TTT ACT TCT TAT ATA GTG TTT GTT ACA Phe Cys Thr Ala His Cys Asn Phe Thr Ser Tyr Ile Val Phe Val Thr 100 105 CAT TGT TTT AAG AGC GGA TCT AAT AGT TGT CCT TTG ACA GGT CTT ATT His Cys Phe Lys Ser Gly Ser Asn Ser Cys Pro Leu Thr Gly Leu Ile CCA AGC GGT TAT ATT CGT ATT GCT GCT ATG AAA CAT GGA AGT CGT ACG 432 Pro Ser Gly Tyr Ile Arg Ile Ala Ala Met Lys His Gly Ser Arg Thr 135 CCT GGT CAC TTA TTT TAT AAC TTA ACA GTT TCT GTG ACT AAA TAT CCT 480 Pro Gly His Leu Phe Tyr Asn Leu Thr Val Ser Val Thr Lys Tyr Pro 150 AAG TTT AGA TCG CTA CAA TGT GTT AAT AAT CAT ACT TCT GTA TAT TTA 528 Lys Phe Arg Ser Leu Gln Cys Val Asn Asn His Thr Ser Val Tyr Leu 170 AAT GGT GAC CTT GTT TTC ACA TCT AAC TAT ACT GAA GAT GTT GTA GCT 576 Asn Gly Asp Leu Val Phe Thr Ser Asn Tyr Thr Glu Asp Val Val Ala 185

GCA GGT GTC CAT TTT AAA AGT GGT GGA CCT ATA ACT TAT AAA GTT ATG

Ala Gly Val His Phe Lys Ser Gly Gly Pro Ile Thr Tyr Lys Val Met

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									GTC Val							672
									GGT Gly							720
AAT Asn	ACT Thr	GGC Gly	AAT Asn	TTT Phe 245	TCA Ser	GAT Asp	GG <b>C</b> Gly	TTC Phe	TAT Tyr 250	CCT Pro	TTT Phe	ACT Thr	AAT Asn	ACT Thr 255	AGT Ser	768
ATT Ile	GTT Val	AAG Lys	GAT Asp 260	AAG Lys	TTT Phe	ATT Ile	GTT Val	TAT Tyr 265	CGT Arg	GAA Glu	AGT Ser	AGT Ser	GTC Val 270	AAT Asn	ACT Thr	816
ACT Thr	TTG Leu	ACA Thr 275	TTA Leu	ACT Thr	AAT Asn	TTC Phe	ACG Thr 280	TTT Phe	AGT Ser	AAT Asn	GAA Glu	AGT Ser 285	GGT Gly	GCC Ala	CCT Pro	864
CCT Pro	AAT Asn 290	ACA Thr	GGT Gly	GGT Gly	GTT Val	GAC Asp 295	AGT Ser	TTT Phe	ATT Ile	TTA Leu	TAC Tyr 300	CAG Gln	ACA Thr	CAA Gln	ACA Thr	912
GCT Ala 305	CAG Gln	AGT Ser	GGT Gly	TAT Tyr	TAT Tyr 310	AAT Asn	TTT Phe	AAT Asn	TTT Phe	TCA Ser 315	TTT Phe	CTG Leu	AGT Ser	AGT Ser	TTT Phe 320	960
GTT Val	TAT Tyr	AGG Arg	GAA Glu	AGT Ser 325	AAT Asn	TAT Tyr	ATG Met	TAT Tyr	GGA Gly 330	TCT Ser	TAC Tyr	CAT His	CCG Pro	GCT Ala 335	TGT Cys	1008
AGT Ser	TTT Phe	AGA Arg	CCT Pro 340	GAA Glu	ACC Thr	CTT Leu	AAT Asn	GGT Gly 345	TTG Leu	TGG Trp	TCT Ser	AAT Asn	TCC Ser 350	CTT Leu	TCT Ser	1056
									GGT Gly							1104
									GCT Ala							1152
	Ala					Tyr			GAG Glu		Thr					1200
									AGC Ser 410							1248
									CAA Gln							1296
									GTT Val							1344
_									GCT Ala							1392
									ACA Thr							1440

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TTC Phe	GTT Val	GTA Val	CAA Gln	GGT Gly 485	GAA Glu	TAT Tyr	GGC Gly	CCT Pro	AAC Asn 490	TAC Tyr	TAT Tyr	AAG Lys	GTT Val	AAT Asn 495	CTA Leu	1488
TGT Cys	GAA Glu	GAT Asp	GTT Val 500	AAC Asn	CAA Gln	CAG Gln	TTT Phe	GTA Val 505	GTT Val	TCT Ser	GGT Gly	GGT Gly	AAA Lys 510	TTA Leu	GTA Val	1536
GGT Gly	ATT Ile	CTC Leu 515	ACT Thr	TCA Ser	CGT Arg	AAT Asn	GAA Glu 520	ACT Thr	GGT Gly	TCT Ser	CAG Gln	CCT Pro 525	CTT Leu	GAA Glu	AAC Asn	1584
														CGT Arg		1632
														AAG Lys		1680
														GAA Glu 575		1728
														CTC Leu		1776
													_	ACG Thr		1824
	GAT Asp 610						A									1846

## (2) INFORMATION FOR SEQ ID NO:17:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 615 amino acids
  - (B) TYPE: amino acid(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Met Leu Val Lys Ser Leu Phe Leu Val Thr Ile Leu Phe Ala Leu Cys

Ser Ala Asn Leu Tyr Asp Asn Glu Ser Phe Val Tyr Tyr Tyr Gln Ser

Ala Phe Arg Pro Gly His Gly Trp His Leu His Gly Gly Ala Tyr Ala

Val Val Asn Val Ser Ser Glu Asn Asn Asn Ala Gly Thr Ala Pro Ser

Cys Thr Ala Gly Ala Ile Gly Tyr Ser Lys Asn Phe Ser Ala Ala Ser

Val Ala Met Thr Ala Pro Leu Ser Gly Met Ser Trp Ser Ala Ser Ser

Phe Cys Thr Ala His Cys Asn Phe Thr Ser Tyr Ile Val Phe Val Thr 105 His Cys Phe Lys Ser Gly Ser Asn Ser Cys Pro Leu Thr Gly Leu Ile Pro Ser Gly Tyr Ile Arg Ile Ala Ala Met Lys His Gly Ser Arg Thr 135 Pro Gly His Leu Phe Tyr Asn Leu Thr Val Ser Val Thr Lys Tyr Pro 155 Lys Phe Arg Ser Leu Gln Cys Val Asn Asn His Thr Ser Val Tyr Leu 170 Asn Gly Asp Leu Val Phe Thr Ser Asn Tyr Thr Glu Asp Val Val Ala Ala Gly Val His Phe Lys Ser Gly Gly Pro Ile Thr Tyr Lys Val Met Arg Glu Val Lys Ala Leu Ala Tyr Phe Val Asn Gly Thr Ala His Asp Val Ile Leu Cys Asp Asp Thr Pro Arg Gly Leu Leu Ala Cys Gln Tyr 230 235 Asn Thr Gly Asn Phe Ser Asp Gly Phe Tyr Pro Phe Thr Asn Thr Ser Ile Val Lys Asp Lys Phe Ile Val Tyr Arg Glu Ser Ser Val Asn Thr Thr Leu Thr Leu Thr Asn Phe Thr Phe Ser Asn Glu Ser Gly Ala Pro 280 Pro Asn Thr Gly Gly Val Asp Ser Phe Ile Leu Tyr Gln Thr Gln Thr Ala Gln Ser Gly Tyr Tyr Asn Phe Asn Phe Ser Phe Leu Ser Ser Phe 315 Val Tyr Arg Glu Ser Asn Tyr Met Tyr Gly Ser Tyr His Pro Ala Cys 325 Ser Phe Arg Pro Glu Thr Leu Asn Gly Leu Trp Ser Asn Ser Leu Ser Val Ser Leu Ile Tyr Gly Pro Ile Gln Gly Gly Cys Lys Gln Ser Val Phe Asn Gly Lys Ala Thr Cys Cys Tyr Ala Tyr Ser Tyr Gly Gly Pro 375 380 Arg Ala Cys Lys Gly Val Tyr Arg Gly Glu Leu Thr Gln His Phe Glu Cys Gly Leu Leu Val Tyr Val Thr Lys Ser Asp Gly Ser Arg Ile Gln Thr Ala Thr Gln Pro Pro Val Leu Thr Gln Asn Phe Tyr Asn Asn Ile 425 Thr Leu Gly Lys Cys Val Asp Tyr Asn Val Tyr Gly Arg Thr Gly Gln

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Gly	Phe 450	Ile	Thr	Asn	Val	Thr 455	Asp	Leu	Ala	Thr	Ser 460	His	Asn	Tyr	Leu
Ala 465	Glu	Gly	Gly	Leu	Ala 470	Ile	Leu	Asp	Thr	Ser 475	Gly	Ala	Ile	Asp	Ile 480
Phe	Val	Val	Gln	Gly 485	Glu	Tyr	Gly	Pro	Asn 490	Tyr	Tyr	Lys	Val	Asn 495	Leu
Суѕ	Glu	Asp	Val 50	Asn O	Gln	Gln	Phe	Val 50	Val	Ser	Gly	Gly	Lys 510		Val
Gly	Ile	Leu 515	Thr	Ser	Arg	Asn	Glu 520	Thr	Gly	Ser	Gln	Pro 525	Leu	Glu	Asn
Gln	Phe 530	Tyr	Ile	Lys	Ile	Thr 535	Asn	Gly	Thr	His	Arg 540	Ser	Arg	Arg	Ser
Val 545	Asn	Glu	Asn	Val	Thr 550	Asn	Cys	Pro	Tyr	Val 555	Ser	Tyr	Gly	Lys	Phe 560
Суѕ	Ile	Lys	Pro	Asp 565	Gly	Ser	Val	Ser	Pro 570	Ile	Val	Pro	Lys	Glu 575	Leu
Glu	Gln	Phe	Val 580	Ala	Pro	Leu	Leu	Asn 585	Val	Thr	Glu	Asn	Val 590	Leu	Ile
Pro	Asn	Ser 595	Phe	Asn	Leu	Thr	Val 600	Thr	Asp	Glu	Tyr	Ile 605	Gln	Thr	Arg
Met	Asp 610	Lys	Val	Gln	Ile	Arg 615									

# (2) INFORMATION FOR SEQ ID NO:18:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2116 base pairs
    (B) TYPE: nucleic acid
    (C) STRANDEDNESS: double
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

TATAATTATC	TAGCAGACGC	AGGTATGGCT	ATTTTAGATA	CATCTGGTTC	CATAGACATC	60
TTTGTTGCAC	AAGGTGAATA	TGGCCTTACT	TATTATAAGG	CTAACCCTTG	CGAAGACGTC	120
AACCAGCAGT	TTGTAGTTTC	TGGTGGTAAA	TTAGTAGGTA	TTCTTACTTC	ACGTAATGAG	180
ACTGGTTCTC	AGCTTCTTGA	GAACCAGTTT	TACATTAAAA	TCACTAATGG	AACACGTCGT	240
TCTAGACGTT	CTATTACTGC	AAATGTHACA	AATYGCCCTT	ATGTTAGCTA	TGGCAAGTTT	300
TGTCTAAAAC	CTGATGGYTC	AGYTTCTGYT	ATAGCACCAC	NNNNNNNNN	מממממממממ	360
иииииииииии	ииииииииии	ממממממממ	ииииииииии	ииииииииии	מממממממממ	420
имимимими	NNNNNNNNN	ממממממממ	NNNNNNNNN	NNNNNNNNN	ТИМИМИМИМ	480
GTTTGTGGCA	ATTCTCTGGA	TTGTAGAAAG	TTGYTTCAAC	AATATGGGCC	TGTTTGBGAC	540

AACATATTGT CTGTGGTAAA	TAGTGTTGGT	CAAAAAGAAG	ATATGGAACT	TCUAAATCTC	600
TATTCTTCTA CTAAACCATC	TGGCTTTAAT	ACACCAGTTT	TTAGTAATCT	YAGCACTGGC	660
GATTTYAATA TTTCTCTTYT	GGTTGACACC	TCCAGTAGTA	CTACTGGGCG	CTCTTTTATT	720
GAAGATCTTT TATTTACAAG	TGTTGAATCT	GTTGGATTAC	CAACAGATGA	AGCTTATAAA	780
AAGTGCACTG CAGGACCTTT	AGGCTTCCTT	AAGGACCTBG	CGTGTGCTCG	TGAATATAAT	840
GGCTTGCTTG YNNNNNNCC	TATTATAACA	GCAGAAATGC	AAACCTTGTA	TACTAGTTCT	900
TTAGTAGCTT CTATGGCTTT	TGGTGGGATT	ACTGCAGCTG	GTGCTATACC	TTTTGCCACA	960
CAACTGCAGG CTAGAATTAA	TCACTTGGGT	ATTACCCAGT	CACTTTTGCA	GAAAAATCAA	1020
GAAAAATTG CTGCCTCCTT	TAATAAGGCC	ATTGGCCATA	TGCAGGAAGG	TTTTAGAAGT	1080
ACATCTCTAG CATTACAACA	AGTYCAMGAT	GTTGTTAATA	AGCAGAGTGC	TATTCTTACT	1140
GAGACTATGG CATCACTTA	TAAAAATTTK	GGTGCTATTT	CTTCTGTGAT	TCAAGATATC	1200
TACCAGCAAC TTGACGCCAT	ACAAGCAAAT	GCTCAAGTGG	ATCGTCTTAT	AACTGGTAGA	1260
TTGTCATCAC TTTCTGTTTT	AGCATCTGCT	AAGCAGGCGG	AGTATATTAG	AGTGTCACAA	1320
CAGCGTGAGT TAGCTACTCA	GAAAATTAAT	GAGTGTGTTA	AATCACAGTC	TATTAGGTAC	1380
TCCTTTTGTG GTAATGGACG	ACACGTTCTA	ACTATACCGC	AAAATGCACC	TAATGGTATA	1440
GTGTTTATAC ACTTTACTTA	TACTCCAGAG	AGTTTTGKTA	ATGTTACTGC	AATAGTGGGT	1500
TTTTGTAARG CCGCTAATGO	TAGTCAGTAT	GCAATAGTGC	CTGCTAATGG	CAGAGGTATT	1560
TCTATACAAG TTAATGGTAG	TCACTACATC	ACTGCACGAG	ATATGTATAT	GCCAAGAGAT	1620
ATTACTGCAG GAGATATAGT	TACGCTTACT	TCTTGTCAAG	CAAATTATGT	AAGTGTAMMT	1680
AAGACCGTCA TTACYACATT	HGTAGACAAT	GATGATTTTG	ATTTTGATGA	CGAATTGTCA	1740
AAATGGTGGA ATGATACTAA	GCATGAGCTA	CCAGACTTTG	ACGAATTCAA	TTACACAGTA	1800
CCTATACTTG ACATTGGTAG	TGAAATTGAT	CGTATTCAAG	GCGTTATACA	GGGCCTTAAT	1860
GACTCTCTAA TAGACCTTGA	AACACTATCA	ATACTCAAAA	CTTATATTAA	GTGGCCTTGG	1920
TATGTGTGGT TAGCCATAGC	TTTTGSCACT	ATTATCTTCA	TCCTAATATT	AGGGTGGGTG	1980
TTTTTCATGA CTGGTTGTTG	TGGTTGTTGT	TGTGGATGCT	TTGGCATTAT	TCCTCTAATG	2040
AGCAAGTGTG GTAAGAAATC	TTCTTATTAC	ACGACTTTGG	ATAATGATGT	GGTAACTGAA	2100
CAAWACAGAC CYAAAA					2116

# (2) INFORMATION FOR SEQ ID NO:19:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 705 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO

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(xi) SE	QUENCE	DESCRIPTION:	SEO	ID	NO:19:
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Tyr Asn Tyr Leu Ala Asp Ala Gly Met Ala Ile Leu Asp Thr Ser Gly Ser Ile Asp Ile Phe Val Ala Gln Gly Glu Tyr Gly Leu Thr Tyr Tyr Lys Ala Asn Pro Cys Glu Asp Val Asn Gln Gln Phe Val Val Ser Gly Gly Lys Leu Val Gly Ile Leu Thr Ser Arg Asn Glu Thr Gly Ser Gln Leu Leu Glu Asn Gln Phe Tyr Ile Lys Ile Thr Asn Gly Thr Arg Arg Ser Arg Arg Ser Ile Thr Ala Asn Val Thr Asn Xaa Pro Tyr Val Ser 90 Tyr Gly Lys Phe Cys Leu Lys Pro Asp Gly Ser Xaa Ser Xaa Ile Ala 120 Val Cys Gly Asn Ser Leu Asp Cys Arg Lys Leu Xaa Gln Gln Tyr Gly Pro Val Xaa Asp Asn Ile Leu Ser Val Val Asn Ser Val Gly Gln Lys Glu Asp Met Glu Leu Leu Asn Leu Tyr Ser Ser Thr Lys Pro Ser Gly Phe Asn Thr Pro Val Phe Ser Asn Leu Ser Thr Gly Asp Phe Asn Ile 215 Ser Leu Leu Val Asp Thr Ser Ser Ser Thr Thr Gly Arg Ser Phe Ile Glu Asp Leu Leu Phe Thr Ser Val Glu Ser Val Gly Leu Pro Thr Asp Glu Ala Tyr Lys Lys Cys Thr Ala Gly Pro Leu Gly Phe Leu Lys Asp 260 Leu Ala Cys Ala Arg Glu Tyr Asn Gly Leu Leu Xaa Xaa Xaa Pro Ile Ile Thr Ala Glu Met Gln Thr Leu Tyr Thr Ser Ser Leu Val Ala Ser 290 Met Ala Phe Gly Gly Ile Thr Ala Ala Gly Ala Ile Pro Phe Ala Thr 310 Gln Leu Gln Ala Arg Ile Asn His Leu Gly Ile Thr Gln Ser Leu Leu Gln Lys Asn Gln Glu Lys Ile Ala Ala Ser Phe Asn Lys Ala Ile Gly 345

His Met Gln Glu Gly Phe Arg Ser Thr Ser Leu Ala Leu Gln Gln Val 360 Xaa Asp Val Val Asn Lys Gln Ser Ala Ile Leu Thr Glu Thr Met Ala Ser Leu Asn Lys Asn Xaa Gly Ala Ile Ser Ser Val Ile Gln Asp Ile 395 Tyr Gln Gln Leu Asp Ala Ile Gln Ala Asn Ala Gln Val Asp Arg Leu Ile Thr Gly Arg Leu Ser Ser Leu Ser Val Leu Ala Ser Ala Lys Gln 425 Ala Glu Tyr Ile Arg Val Ser Gln Gln Arg Glu Leu Ala Thr Gln Lys Ile Asn Glu Cys Val Lys Ser Gln Ser Ile Arg Tyr Ser Phe Cys Gly Asn Gly Arg His Val Leu Thr Ile Pro Gln Asn Ala Pro Asn Gly Ile Val Phe Ile His Phe Thr Tyr Thr Pro Glu Ser Phe Xaa Asn Val Thr 485 490 Ala Ile Val Gly Phe Cys Lys Ala Ala Asn Ala Ser Gln Tyr Ala Ile 505 Val Pro Ala Asn Gly Arg Gly Ile Ser Ile Gln Val Asn Gly Ser His 515 Tyr Ile Thr Ala Arg Asp Met Tyr Met Pro Arg Asp Ile Thr Ala Gly 535 Asp Ile Val Thr Leu Thr Ser Cys Gln Ala Asn Tyr Val Ser Val Xaa Lys Thr Val Ile Thr Thr Xaa Val Asp Asp Asp Phe Asp Phe Asp Asp Glu Leu Ser Lys Trp Trp Asn Asp Thr Lys His Glu Leu Pro Asp 585 Phe Asp Glu Phe Asn Tyr Thr Val Pro Ile Leu Asp Ile Gly Ser Glu Ile Asp Arg Ile Gln Gly Val Ile Gln Gly Leu Asn Asp Ser Leu Ile Asp Leu Glu Thr Leu Ser Ile Leu Lys Thr Tyr Ile Lys Trp Pro Trp 630 635 Tyr Val Trp Leu Ala Ile Ala Phe Xaa Thr Ile Ile Phe Ile Leu Ile Leu Gly Trp Val Phe Phe Met Thr Gly Cys Cys Gly Cys Cys Gly Cys Phe Gly Ile Ile Pro Leu Met Ser Lys Cys Gly Lys Lys Ser Ser 680 Tyr Tyr Thr Thr Leu Asp Asn Asp Val Val Thr Glu Gln Xaa Arg Pro 695

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Tys 705	
(2) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 36 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
GAATTCGAGC TCGCCCGGGG ATCCTCTAGA GTCGAC	36
(2) INFORMATION FOR SEQ ID NO:21:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 57 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1357	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CACAGCTCAA CA ATG AAG TGG GCA ACG TGG ATC GAT CCC GTC GTT TTA  Met Lys Trp Ala Thr Trp Ile Asp Pro Val Val Leu  1 5 10	48
CAA CGT CGT Gln Arg Arg 15	57
(2) INFORMATION FOR SEQ ID NO:22:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 15 amino acids</li> <li>(B) TYPE: amino acid</li> <li>(D) TOPOLOGY: linear</li> </ul>	

Met Lys Trp Ala Thr Trp Ile Asp Pro Val Val Leu Gln Arg Arg 1 5 10 10

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

(ii) MOLECULE TYPE: protein

(2) INF	FORMATION FOR SEQ ID NO:23:	
(i	(A) LENGTH: 57 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
(ii	) MOLECULE TYPE: DNA (genomic)	
(iii	) HYPOTHETICAL: NO	
(iv	) ANTI-SENSE: NO	
(xi	) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
ACTCGGG	CAG CGTTGGGTCC TGGGACTCTA GAGGATCGAT CCCCTATGGC GATCATC	57
(2) INF	ORMATION FOR SEQ ID NO:24:	
(i	) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 99 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
(ii	) MOLECULE TYPE: DNA (genomic)	
(iii	) HYPOTHETICAL: NO	
(iv	) ANTI-SENSE: NO	
(xi	) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
GCGCCCA	CGT GGCCTGGTAC AATTCGAGCT CGCCCGGGGA TCCTCTAGAG TCGACTCTAG	60
AGGATCG	ATC CTCTAGAGTC GGCGGGACGA GCCCGCGAT	99
(2) INF	ORMATION FOR SEQ ID NO:25:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 57 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:25:	
TCCACAGG	AC CTGCAGCGAC CCGCTTAACA GCGTCAACAG CGTGCCGCAG ATCGGGG	57
(2) INFO	RMATION FOR SEQ ID NO:26:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 35 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	

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(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:26:	
GTTGATCC	CG GGAGATGGGG GAGGCTAACT GAAAC	35
(2) INFO	RMATION FOR SEQ ID NO:27:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 103 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:27:	
GCTCATGG	TG GCCCCGGGC GGTTCAACGA GGGCCAGTAC CGGCGCCTGG TGTCCGTCGA	60
CCTGCAGG	IC GACTCTAGAG GATCCCCGGG CGAGCTCGAA TTC	103
(2) INFO	RMATION FOR SEQ ID NO:28:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 66 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:28:	
GAATTCGA(	GC TCGCCCGGGG ATCCTCTAGA GTCGACGTCT GGGGCGCGGG GGTGGTGCTC	60
TTCGAG		66
(2) INFO	RMATION FOR SEQ ID NO:29:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 66 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	

(iv) ANTI-SENSE: NO

(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1666	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
CTCCACAGCT CAACA ATG AAG TGG GCA ACG TGG ATC GAT CCC GTC GTT TTA  Met Lys Trp Ala Thr Trp Ile Asp Pro Val Val Leu  1 5 10	51
CAA CGT CGT GAC TGG Gln Arg Arg Asp Trp 15	66
(2) INFORMATION FOR SEQ ID NO:30:	
<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 17 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>	
(ii) MOLECULE TYPE: protein	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
Met Lys Trp Ala Thr Trp Ile Asp Pro Val Val Leu Gln Arg Arg Asp 1 5 10 15	•
Trp	
(2) INFORMATION FOR SEQ ID NO:31:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 132 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 193	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
GAC GAC TCC TGG AGC CCG TCA GTA TCG GCG GAA ATC CAG CTG AGC GCC Asp Asp Ser Trp Ser Pro Ser Val Ser Ala Glu Ile Gln Leu Ser Ala 1 5 10	48
GGT CGC TAC CAT TAC CAG TTG GTC TGG TGT CAA AAA GAT CTA GAA Gly Arg Tyr His Tyr Gln Leu Val Trp Cys Gln Lys Asp Leu Glu 20 25 30	93
TAAGCTAGAG GATCGATCCC CTATGGCGAT CATCAGGGC	132
(2) INFORMATION FOR SEQ ID NO:32:	

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 31 amino acids

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(B) TYPE: amino acid (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: protein	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
Asp Asp Ser Trp Ser Pro Ser Val Ser Ala Glu Ile Gln Leu Ser Ala 1 5 10 15	
Gly Arg Tyr His Tyr Gln Leu Val Trp Cys Gln Lys Asp Leu Glu 20 25 30	
(2) INFORMATION FOR SEQ ID NO:33:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 66 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
AACGAGGGCC AGTACCGGCG CCTGGTGTCC GTCGACTCTA GAGGATCCCC GGGCGAGCTC	60
GAATTC	66
(2) INFORMATION FOR GEO ID NO. 24.	
(2) INFORMATION FOR SEQ ID NO:34:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 65 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
CAGGTCGAAG CTTGGGCGCT GCCTATGTAG TGAAATCTAT ACTGGGATTT ATCATAACTA	60
GTTTA	65
(2) INFORMATION FOR SEQ ID NO:35:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 65 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	

(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:	
AATAATCTAT CACTTTGTCA TGGAGATGCC CAAGCTTCGA CGACTCCCTT GGCCATGATG	60
AATGG	65
(2) INFORMATION FOR SEQ ID NO:36:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 65 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
TATACCAGCT ACGGCGCTAG CATTCATGGT ATCCCGTGAT TGCTCGATGC TTTCCTTCTG	60
AATTC	65 •
(2) INFORMATION FOR SEQ ID NO:37:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 65 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:	
AAGCTTGGCC TCGTCGTTAA TTAACCCAAT TCGAGCTCGC CCAGCTTGGG CTGCAGGTCG	60
GGAAC	65
(2) INFORMATION FOR SEQ ID NO:38:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 65 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION, SEC ID NO. 20.	

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TGTTTCAGTT AGCCTCCCCC ATCTCCCGAC TCTAGAGGAT CTCGACATAG CGAATACATT	60
TATGG	65
(2) INFORMATION FOR SEQ ID NO:39:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 130 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:	
AACGTATATA TTTTTCACGA CGTAGACCAC TATTGCCATG GACTCTAGAG GATCGGGTAC	60
CGAGCTCGAA TTGGGAAGCT TGTCGACTTA ATTAAGCGGC CGCGTTTAAA CGGCCCTCGA	120
GGCCAAGCTT	130
(2) INFORMATION FOR SEQ ID NO:40:	•
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 60 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:	
GTCGACGTCT GGGGCGCGGG GGTGGTGCTC TTCGAGACGC TGCCTACCCC AAGACGATCG	60
(2) INFORMATION FOR SEQ ID NO:41:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 60 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:	
AGCTCAACAA TGAAGTGGGC AACGTGGATC GATCCCGTCG TTTTACAACG TCGTGACTGG	60

(2) INFORMATION FOR SEQ ID NO:42:

(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 120 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:	
GAGCCCGTCA GTATCGGCGG AAATCCAGCT GAGCGCCGGT CGCTACCA	TT ACCAGTTGGT 60
GTTGGTCTGG TGTCAAAAAG ATCCGGACCG CGCCGTTAGC CAAGTTGC	GT TAGAGAATGA 120
(2) INFORMATION FOR SEQ ID NO:43:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 60 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: DNA (genomic)	•
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
ACACAGTCAC ACTCATGGGG GCCGAAGGCA GAATTCGTAA TCATGGTCA	AT AGCTGTTTCC 60
(2) INFORMATION FOR SEQ ID NO:44:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 60 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
AAACCTGTCG TGCCAGCGAG CTCGGGATCC TCTAGAGGAT CCCCGGGCC	CC CGCCCCCTGC 60
(2) INFORMATION FOR SEQ ID NO:45:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 60 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (GODONIA)	

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(iii) HYPOTHETICAL: NO
     (iv) ANTI-SENSE: NO
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:
TCGTCCACAC GGAGCGCGGC TGCCGACACG GATCCCGGTT GGCGCCCTCC AGGTGCAGGA
                                                                        60
(2) INFORMATION FOR SEO ID NO:46:
     (i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 60 base pairs
           (B) TYPE: nucleic acid
           (C) STRANDEDNESS: double
           (D) TOPOLOGY: linear
    (ii) MOLECULE TYPE: DNA (genomic)
   (iii) HYPOTHETICAL: NO
    (iv) ANTI-SENSE: NO
    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:
AACCCCCCC CCCCCCCCC CCCCCCCTG CAGGCATCGT GGTGTCACGC TCGTCGTTTG
                                                                        60
(2) INFORMATION FOR SEQ ID NO:47:
     (i) SEQUENCE CHARACTERISTICS:
          (A) LENGTH: 60 base pairs
          (B) TYPE: nucleic acid
          (C) STRANDEDNESS: double
          (D) TOPOLOGY: linear
    (ii) MOLECULE TYPE: DNA (genomic)
   (iii) HYPOTHETICAL: NO
    (iv) ANTI-SENSE: NO
    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:
TGTCATGCCA TCCGTAAGAT GCTTTTCTGT GACTGGTGAG TCGGATCCTC TAGAGTCGAC
                                                                        60
(2) INFORMATION FOR SEQ ID NO:48:
     (i) SEQUENCE CHARACTERISTICS:
          (A) LENGTH: 2681 base pairs
          (B) TYPE: nucleic acid
          (C) STRANDEDNESS: double
          (D) TOPOLOGY: linear
    (ii) MOLECULE TYPE: DNA (genomic)
   (iii) HYPOTHETICAL: NO
    (iv) ANTI-SENSE: NO
    (ix) FEATURE:
          (A) NAME/KEY: CDS
          (B) LOCATION: 146..481
    (ix) FEATURE:
```

(A) NAME/KEY: CDS

(B) LOCATION: complement (602..1402)

# (ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1599..2135

## (ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: complement (2308..2634)

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:

TTTATCGGAC	CTTGGGTATT	CAGGGGAACC	CATCTGGTTG	AAATGCATCC	GACCCTGCAC	60
TTGATCCTGG	TTACCCCGAC	CCAANTTTTA	AGCCGGCTGG	CGCGGTCCCT	AGATAACCCC	120
CCGCTTAAAA	CTAGCCCCAA	TATTGATGTG	CAGATATAAC	ACAGNNANCC	GATCAATGGA	180
AGACATGCTA	CGGCGGTCAT	CTCCCGAAGA	CATCACCGAT	TCCCTAACAA	TGTGCCTGAT	240
TATGTTATCG	CGCATTCGTC	GTACCATGCG	CACCGCAGGA	AATAAATATA	GCTATATGAT	300
AGATCCAATG	AATCGTATGT	CTAATTACAC	TCCAGGCGAA	TGTATGACAG	GTATATTGCG	360
ATATATTGAC	GAACATGCTA	GAAGGTGTCC	TGATCACATA	TGTAATTTGT	ATATCACATG	420
TACACTTATG	CCGATGTATG	TGCACGGGCG	ATATTTCTAT	TGTAATTCAT	TTTTTTGKTA	480
GTAAACTACC	ACAGGCTGTC	CGGAAATCTA	AGTTAATGAA	TAAAGTAGAT	GGTTAATACT	540
CATTGCTTAG	AATTGGACTA	CTTTTAATYC	TCTTTAATGT	TCGTATTAAA	TAAAAACATC	600
TTTAATAAAC	TTCAGCCTCT	TCGCTTATTG	TAGAAATTGA	GTATTCAMAA	TCATGTTCAA	660
AGCCGTCTTC	GGAGAGTGTA	CTCGCCACGG	TGGTTGGAAC	ATCACTATGT	CTACACGTCA	720
AATTTAAGCA	CGTCAGGTCT	GTCGAGGACA	AGAAATGGTT	AACTAGTGTT	TCAATTATTC	780
TTATAAACGT	TAAGCATTGT	AAGCCCCCCG	GCCGTCCGCA	GCAACAATTT	ACTAGTATGC	840
CGTGGGCTCC	GGGACTATCA	CGGATGTCCA	ATTCGCACAT	GCATATAATT	TTTCTAGGGT	900
CTCTCATTTC	GAGAAATCTT	CGGGGATCCA	TCAGCAATGC	GGGCTGTAGT	CCCGATTCCC	960
GTTTCAAATG	AAGGTGCTCC	AACACGGTCT	TCAAAGCAAC	CGGCATACCA	GCAAACACAG	1020
ACTGCAACTC	CCCGCTGCAA	TGATTGGTTA	TAAACAGTAA	TCTGTCTTCT	GGAAGTATAT	1080
TTCGCCCGAC	AATCCACGGC	GCCCCCAAAG	TTAAAAACCA	TCCATGTGTA	TTTGCGTCTT	1140
CTCTGTTAAA	AGAATATTGA	CTGGCATTTT	CCCGTTGACC	GCCAGATATC	CAAAGTACAG	1200
CACGATGTTG	CACGGACGAC	TTTGCAGTCA	CCAGCCTTCC	TTTCCACCCC	CCCACCAACA	1260
AAATGTTTAT	CGTAGGACCC	ATATCCGTAA	TAAGGATGGG	TCTGGCAGCA	ACCCCATAGG	1320
CGCCTCGGCG	TGGTAGTTCT	CGAGGATACA	TCCAAAGAGG	TTGAGTATTC	TCTCTACACT	1380
TCTTGTTAAA	TGGAAAGTGC	ATTTGCTTGT	TCTTACAATC	GGCCCGAGTC	TCGTTCACAG	1440
CGCCTCGTTC	ACACTTAAAC	CACAAATAGT	CTACAGGCTA	TATGGGAGCC	AGACTGAAAC	1500
TCACATATGA	CTAATATTCG	GGGGTGTTAG	TCACGTGTAG	CCCATTGTGT	GCATATAACG	1560
ATGTTGGACG	CGTCCTTATT	CGCGGTGTAC	TTGATACTAT	GGCAGCGAGC	ATGGGATATT	1620
CATCCTCGTC	ATCGTTAACA	TCTCTACGGG	TTCAGAATGT	TTGGCATGTC	GTCGATCCTT	1680
TGCCCATCGT	TGCAAATTAC	AAGTCCGATC	GCCATGACCG	CGATAAGCCT	GTACCATGTG	1740

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GCATTAGGGT	GACATCTCGA	TCATACATTA	TAAGACCAAC	GTGCGAGTCT	TCCAAAGACC	1800
TGCACGCCTT	CTTCTTCGGA	TTGTCAACGG	GTTCTTCAGA	ATCTATGCCC	ATATCTGGCG	1860
TTGAGACCAT	TGTGCGTTTA	ATGAACAATA	AAGCGGCATG	CCATGGAAAG	GAGGCTGCA	1920
GATCTCCATT	TTCTCACGCC	ACTATCCTGG	ACGCTGTAGA	CGATAATTAT	ACCATGAATA	1980
TAGAGGGGGT	ATGTTTCCAC	TGCCACTGTG	ATGATAAGTT	TTCTCCAGAT	TGTTGGATAT	2040
CTGCATTTTC	TGCTGCCGAA	CAAACTTCAT	CGCTATGCAA	AGAGATGCGT	GTGTACACGC	2100
GCCGGTGGAG	TATACGGGAA	ACTAAATGTT	CATAGAGGTC	TTTGGGCTAT	ATGTTATTAA	2160
TAATAAATA	TGACCAGTGA	ACAATTTGTT	TAATGTTAGT	TTATTCAATG	CATTGGTTGC	2220
AAATATTCAT	TACTTCTCCA	ATCCCAGGTC	ATTCTTTAGC	GAGATGATGT	TATGACATTG	2280
CTGTGAAAAT	TACTACAGGA	TATATTTTTA	AGATGCAGGA	GTAACAATGT	GCATAGTAGG	2340
CGTAGTTATC	GCAGACGTGC	AACGCTTCGC	ATTTGAGTTA	CCGAAGTGCC	CAACAGTGCT	2400
GCGGTTATGG	TTTATGCGCA	CAGAATCCAT	GCATGTCCTA	ATTGAACCAT	CCGATTTTTC	2460
TTTTAATCGC	GATCGATGTT	TGGGCAACTG	CGTTATTTCA	GATCTAAAAA	ATTTACCCTY	2520
TATGACCATC	ACATCTCTCT	GGYTCATACC	CCGCTTGGGN	TAAGATATCA	TGTAGATTCC	2580
GCCCCTAAGA	AATTGCAAAC	TAACATNATT	GNCGGGTTCC	ATATACAATC	CCATCTTGTC	2640
CNCTCGAAAT	TACAAACTCG	CGCAATAGAC	CCCCGTACAT	T		2681

# (2) INFORMATION FOR SEQ ID NO:49:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 111 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

Met Cys Arg Tyr Asn Thr Xaa Xaa Arg Ser Met Glu Asp Met Leu Arg 1 5 10 15

Arg Ser Ser Pro Glu Asp Ile Thr Asp Ser Leu Thr Met Cys Leu Ile
20 25 30

Met Leu Ser Arg Ile Arg Arg Thr Met Arg Thr Ala Gly Asn Lys Tyr

Ser Tyr Met Ile Asp Pro Met Asn Arg Met Ser Asn Tyr Thr Pro Gly 50 55 60

Glu Cys Met Thr Gly Ile Leu Arg Tyr Ile Asp Glu His Ala Arg Arg 65 70 75 80

Cys Pro Asp His Ile Cys Asn Leu Tyr Ile Thr Cys Thr Leu Met Pro

Met Tyr Val His Gly Arg Tyr Phe Tyr Cys Asn Ser Phe Phe Xaa 100 105 110

## (2) INFORMATION FOR SEQ ID NO:50:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 266 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

Met His Phe Pro Phe Asn Lys Lys Cys Arg Glu Asn Thr Gln Pro Leu 1 5 10 15

Trp Met Tyr Pro Arg Glu Leu Pro Arg Arg Gly Ala Tyr Gly Val Ala
20 25 30

Ala Arg Pro Ile Leu Ile Thr Asp Met Gly Pro Thr Ile Asn Ile Leu
35 40 • 45

Leu Val Gly Gly Trp Lys Gly Arg Leu Val Thr Ala Lys Ser Ser Val
50 60

Gln His Arg Ala Val Leu Trp Ile Ser Gly Gly Gln Arg Glu Asn Ala 65 70 75 80

Ser Gln Tyr Ser Phe Asn Arg Glu Asp Ala Asn Thr His Gly Trp Phe 85 90 95

Leu Thr Leu Gly Ala Pro Trp Ile Val Gly Arg Asn Ile Leu Pro Glu 100 105 110

Asp Arg Leu Leu Phe Ile Thr Asn His Cys Ser Gly Glu Leu Gln Ser 115 120 125

Val Phe Ala Gly Met Pro Val Ala Leu Lys Thr Val Leu Glu His Leu 130 135 140

His Leu Lys Arg Glu Ser Gly Leu Gln Pro Ala Leu Leu Met Asp Pro 145 150 155 160

Arg Arg Phe Leu Glu Met Arg Asp Pro Arg Lys Ile Ile Cys Met Cys 165 170 175

Glu Leu Asp Ile Arg Asp Ser Pro Gly Ala His Gly Ile Leu Val Asn 180 185 190

Cys Cys Cys Gly Arg Pro Gly Gly Leu Gln Cys Leu Thr Phe Ile Arg 195 200 205

Ile Ile Glu Thr Leu Val Asn His Phe Leu Ser Ser Thr Asp Leu Thr 210 215 220

Cys Leu Asn Leu Thr Cys Arg His Ser Asp Val Pro Thr Thr Val Ala 225 230 235 240

Ser Thr Leu Ser Glu Asp Gly Phe Glu His Asp Xaa Glu Tyr Ser Ile 245 250 255 209

Ser Thr Ile Ser Glu Glu Ala Glu Val Tyr 260 265

- (2) INFORMATION FOR SEQ ID NO:51:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 178 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: double
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

Met Ala Ala Ser Met Gly Tyr Ser Ser Ser Ser Leu Thr Ser Leu

1 10 15

Arg Val Gln Asn Val Trp His Val Val Asp Pro Leu Pro Ile Val Ala 20 25 30

Asn Tyr Lys Ser Asp Arg His Asp Arg Asp Lys Pro Val Pro Cys Gly 35 40 . 45

Ile Arg Val Thr Ser Arg Ser Tyr Ile Ile Arg Pro Thr Cys Glu Ser 50 55 60

Ser Lys Asp Leu His Ala Phe Phe Phe Gly Leu Ser Thr Gly Ser Ser 65 70 75 80

Glu Ser Met Pro Ile Ser Gly Val Glu Thr Ile Val Arg Leu Met Asn 85 90 95

Asn Lys Ala Ala Cys His Gly Lys Glu Gly Cys Arg Ser Pro Phe Ser

His Ala Thr Ile Leu Asp Ala Val Asp Asp Asn Tyr Thr Met Asn Ile 115 120 125

Glu Gly Val Cys Phe His Cys His Cys Asp Asp Lys Phe Ser Pro Asp

Cys Trp Ile Ser Ala Phe Ser Ala Ala Glu Gln Thr Ser Ser Leu Cys 145 150 155 160

Lys Glu Met Arg Val Tyr Thr Arg Arg Trp Ser Ile Arg Glu Thr Lys 165 170 175

Cys Ser

- (2) INFORMATION FOR SEQ ID NO:52:
  - (i) SEOUENCE CHARACTERISTICS:
    - (A) LENGTH: 108 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: double
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO

- (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

Met Gly Leu Tyr Met Glu Pro Xaa Asn Xaa Val Ser Leu Gln Phe Leu

Arg Gly Gly Ile Tyr Met Ile Ser Xaa Pro Lys Arg Gly Met Xaa Gln

Arg Asp Val Met Val Ile Xaa Gly Lys Phe Phe Arg Ser Glu Ile Thr

Gln Leu Pro Lys His Arg Ser Arg Leu Lys Glu Lys Ser Asp Gly Ser

Ile Arg Thr Cys Met Asp Ser Val Arg Ile Asn His Asn Arg Ser Thr

Val Gly His Phe Gly Asn Ser Asn Ala Lys Arg Cys Thr Ser Ala Ile

Thr Thr Pro Thr Met His Ile Val Thr Pro Ala Ser 100

- (2) INFORMATION FOR SEQ ID NO:53:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 37 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA Oligonucleotide Primer
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: YES
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

# CTCGCTCGCC CATGATCATT AAGCAAGAAT TCCGTCG

37

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- (2) INFORMATION FOR SEQ ID NO:54:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid

    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA Oligonucleotide Primer
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:

# CTGGTTCGGC CCATGATCAG ATGACAAACC TGCAAGATC

- (2) INFORMATION FOR SEQ ID NO:55:
  - (i) SEQUENCE CHARACTERISTICS:

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		<ul><li>(A) LENGTH: 57 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: double</li><li>(D) TOPOLOGY: linear</li></ul>	
	(ii)	MOLECULE TYPE: DNA (genomic)	
(	(iii)	HYPOTHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:55:	
CTCG	GCGT	GG TAGTTCTCGA GGCCTTAATT AAGGCCCTCG AGGATACATC CAAAGAG	51
(2)	INFO	RMATION FOR SEQ ID NO:56:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 63 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
(	(iii)	HYPOTHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:56:	
CGGC	GTGG	TA GTTCTCGAGG CCTTAAGCGG CCGCTTAAGG CCCTCGAGGA TACATCCAAA	60
GAG			63
(2)	TNFO	RMATION FOR SEQ ID NO:57:	
()		SEQUENCE CHARACTERISTICS:  (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
(	(iii)	HYPOTHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:57:	
CGCA	AGGAT	CC GGGGCGTCAG AGGCGGGCGA GGTG	34
(2)	INFO	RMATION FOR SEQ ID NO:58:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 32 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	

(iii) HYPOTHETICAL: NO

(1V)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:58:	
GAGCGGAT	CC TGCAGGAGGA GACACAGAGC TG	32
(2) INFO	RMATION FOR SEQ ID NO:59:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 33 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:59:	
TGTAGAGA'	IC TGGCTAAGTG CGCGTGTTGC CTG	33
(2) INFO	RMATION FOR SEQ ID NO:60:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 33 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:60:	
TGTACAGA:	TC TCACCATGGC TGTGCCTGCA AGC	33

# What is claimed is:

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- 1. A recombinant herpesvirus of turkeys comprising a foreign DNA sequence encoding a cytokine inserted into an insertion region which comprises a XhoI site within a EcoR1 #9 fragment of a herpesvirus of turkeys viral genome, and the foreign DNA sequence encoding a cytokine which is capable of being expressed in a host cell infected with the herpesvirus of turkeys.
- 10 2. The recombinant herpesvirus of turkeys of claim 1, wherein the cytokine is chicken myelomonocytic growth factor (cMGF), chicken interferon (cIFN), interleukin-2, interleukin-6, interleukin-12, interferons, granulocyte-macrophage colony stimulating factors, or interleukin receptors.
  - The recombinant herpesvirus of turkeys of claim
     further comprising a second foreign DNA sequence.
  - The recombinant herpesvirus of turkeys of claim
     wherein the foreign DNA sequence encodes a polypeptide.
- 5. The recombinant herpesvirus of turkeys of claim4, wherein the polypeptide is antigenic.
- 6. The recombinant herpesvirus of turkeys of claim 4, wherein the polypeptide is *E. coli* beta-30 galactosidase.
  - 7. The recombinant herpesvirus of turkeys of claim 2, which is designated S-HVT-144.

- 8. The recombinant herpesvirus of turkeys of claim 5, wherein the foreign DNA sequence encoding an antigenic polypeptide is inserted into an insertion region of the herpesvirus of turkeys viral genome comprising a unique StuI site within the US2 gene.
- 9. The recombinant herpesvirus of turkeys of claim 8, wherein the foreign DNA sequence encodes an 10 antigenic polypeptide selected from the group consisting of: Marek's disease virus, Newcastle disease virus, Infectious laryngotracheitis virus. Infectious bronchitis virus, and Infectious bursal disease virus.

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10. The recombinant herpesvirus of turkeys of claim 9, wherein the foreign DNA sequence encodes Marek's disease virus glycoprotein A, Marek's

disease virus glycoprotein B or Marek's disease virus glycoprotein D.

- 11. The recombinant herpesvirus of turkeys of claim 9, wherein the foreign DNA sequence encodes Newcastle disease virus fusion protein or Newcastle disease virus hemagglutinin-neuraminidase.
- 12. The recombinant herpesvirus of turkeys of claim 9, wherein the foreign DNA sequence encodes Infectious laryngotracheitis virus glycoprotein B, Infectious laryngotracheitis virus glycoprotein I or Infectious laryngotracheitis virus glycoprotein D.
- 35 13. The recombinant herpesvirus of turkeys of claim 9, wherein the foreign DNA sequence encodes

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Infectious bronchitis virus spike protein or Infectious bronchitis virus matrix protein.

- The recombinant herpesvirus of turkeys of claim
  9, wherein the foreign DNA sequence encodes
  Infectious bursal disease virus VP2, Infectious
  bursal disease virus VP3, or Infectious bursal
  disease virus VP4.
- 15. The recombinant herpesvirus of turkeys of claim 1, wherein the cytokine is under control of an endogenous upstream herpesvirus promoter.
- 16. The recombinant herpesvirus of turkeys of claim
  15. the recombinant herpesvirus of turkeys of claim
  15. the recombinant herpesvirus of turkeys of claim
  16. herein the cytokine is under control of a
  16. heterologous upstream promoter.
- 17. The recombinant herpesvirus of turkeys of claim 15, wherein the promoter is selected from PRV gX, HSV-1 alpha 4, HCMV immediate early, MDV gA, MDV gB, MDV gD, ILT gB, BHV-1.1 VP8 and ILT gD.
- 18. A homology vector for producing a recombinant herpesvirus of turkeys by inserting a foreign DNA sequence encoding a cytokine into the viral genome of a herpesvirus of turkey which comprises a double-stranded DNA molecule consisting essentially of:
- a) double stranded foreign DNA not usually present within the herpesvirus of turkeys viral genome;

- b) at one end the foreign DNA, doublestranded herpesvirus of turkeys DNA homologous to the viral genome located at one side of the EcoR1 #9 fragment of the coding region of the herpesvirus of turkeys viral genome; and
- at the other end of the foreign DNA, double-stranded herpesvirus of turkeys

  DNA homologous to the viral genome located at the other side of the EcoR1 #9 of the coding region of the herpesvirus of turkeys viral genome.

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- 15 19. The recombinant herpesvirus of turkeys of claim 18, the cytokine is chicken wherein myelomonocytic growth factor (cMGF), chicken interferon (cIFN), interleukin-2, interleukin-6, interleukin-12, interferons. granulocytemacrophage colony stimulating 20 factors, interleukin receptors.
- 20. A homology vector of claim 18, further comprising
  a second foreign DNA sequence encoding an
  antigenic polypeptide
  - 21. A homology vector of claim 20, wherein the antigenic polypeptide is selected from a group consisting essentially of: Marek's disease virus, Newcastle disease virus, Infectious laryngotracheitis virus, Infectious bronchitis virus and Infectious bursal disease virus.
- 35 22. A homology vector of claim 20, wherein the antigenic polypeptide is selected from a group consisting essentially of: Marek's disease virus

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glycoprotein Α, Marek's disease virus glycoprotein B, Marek's disease virus glycoprotein D, Newcastle disease virus fusion protein, Newcastle disease virus hemagglutininneuraminidase, Infectious laryngotracheitis virus glycoprotein B, Infectious laryngotracheitis virus glycoprotein I, Infectious laryngotracheitis virus glycoprotein Infectious bronchitis virus spike protein, Infectious bronchitis virus matrix protein, Infectious bursal disease virus VP2, Infectious bursal disease virus VP3, and Infectious bursal disease virus VP4.

- 15 23. The homology vector of claim 20, wherein the foreign DNA sequence encodes a screenable marker.
- 24. The homology vector of claim 23, wherein the screenable marker is *E. coli B*-galactosidase or *E. coli B*-glucuronidase.
  - 25. The homology vector of claim 18 designated 751-87.A8.
- 25 26. The homology vector of claim 18 designated 761-07.A1.
- A vaccine useful for immunizing a bird against
  Marek's disease virus which comprises an
  effective immunizing amount of the recombinant
  herpesvirus of turkeys of claims 10 and a
  suitable carrier.
- 28. A vaccine useful for immunizing a bird against

  Newcastle disease virus virus which comprises an
  effective immunizing amount of the recombinant

herpesvirus of turkeys of claim 11 and a suitable carrier.

- 29. A vaccine useful for immunizing a bird against Infectious laryngotracheitis virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys of claim 12 and a suitable carrier.
- 30. A multivalent vaccine useful for immunizing a bird against Marek's disease virus and Newcastle disease virus which comprises an effective immunizing amount of the recombinant herpesvirus of turkeys of claims 11.
- 31. A method of immunizing a bird against Marek's disease virus which comprises administering to the bird an effecting immunizing dose of the vaccine of claim 27.
- 32. A host cell infected with the recombinant herpesvirus of turkey of claim 1.

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- 33. A host cell of claim 32, wherein the host cell is an avian cell.
- 34. A recombinant herpesvirus of turkeys-Marek's disease virus chimera comprising a herpesvirus of turkeys unique long viral genome region and a Marek's disease virus unique short region.
  - 35. The recombinant herpesvirus of turkeys-Marek's disease virus chimera of claim 34, wherein a foreign DNA sequence is inserted within the EcoRl #9 fragment of the herpesvirus of turkeys viral genome, and is capable of being expressed in a

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host cell infected with the herpesvirus of turkeys.

- 36. The recombinant herpesvirus of turkeys-Marek's disease virus chimera of claim 35, wherein the foreign DNA sequence encodes a polypeptide.
- 37. The recombinant herpesvirus of turkeys-Marek's disease virus chimera of claim 36, wherein the foreign DNA sequence encodes a cytokine.

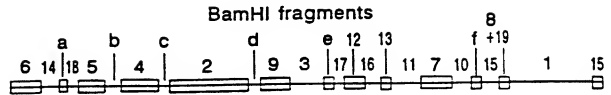
15

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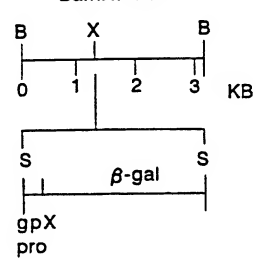
- 38. The recombinant herpesvirus of turkeys-Marek's disease virus chimera of claim 37, wherein the cytokine is a chicken mylomonocytic growth factor (cMGF) or chicken interferon (cIFN).
- 39. The recombinant herpesvirus of turkeys-Marek's disease virus chimera of claim 38, further comprising a foreign DNA sequence encoding the antigenic polypeptide selected from the group consisting of: Marek's disease virus, Newcastle disease virus, Infectious laryngotracheitis virus, Infectious bronchitis virus and Infectious bursal disease virus.
- The recombinant herpesvirus of turkeys of claim 39, designated S-HVT-145.

## FIGURE 1A

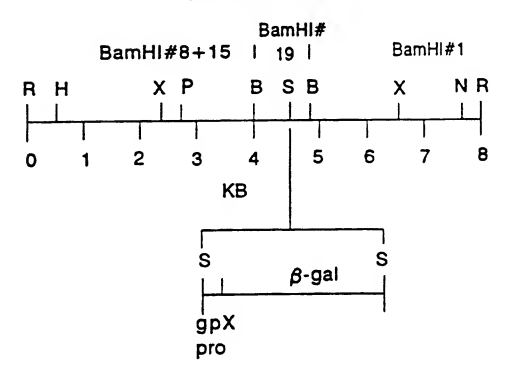


## FIGURE 1B

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## FIGURE 1C



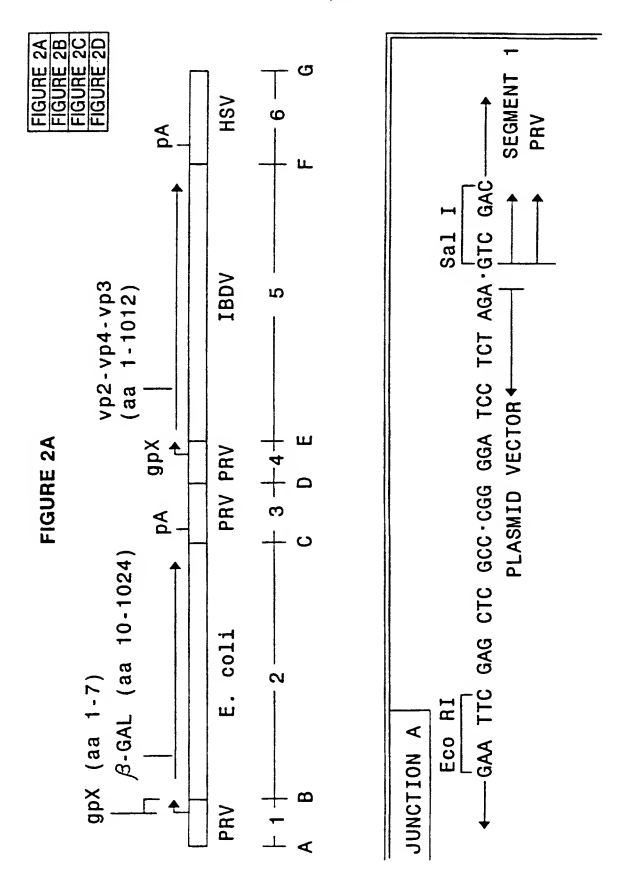
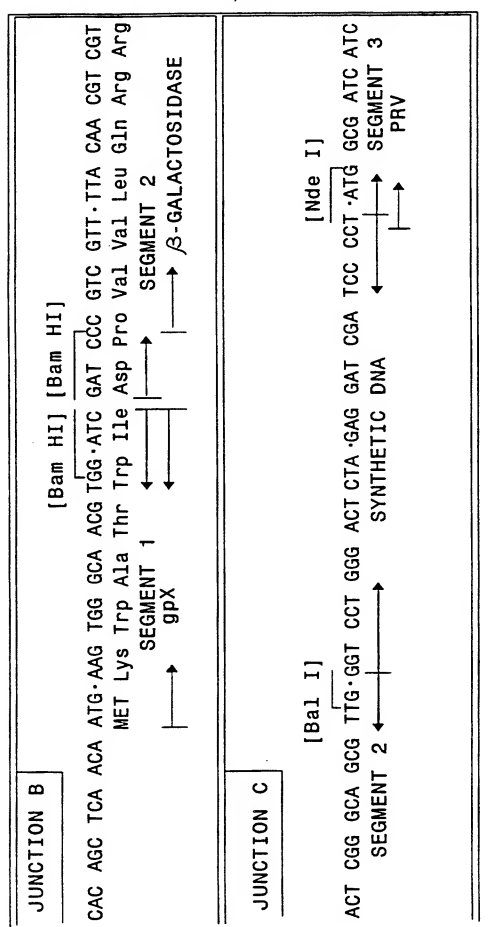


FIGURE 2B



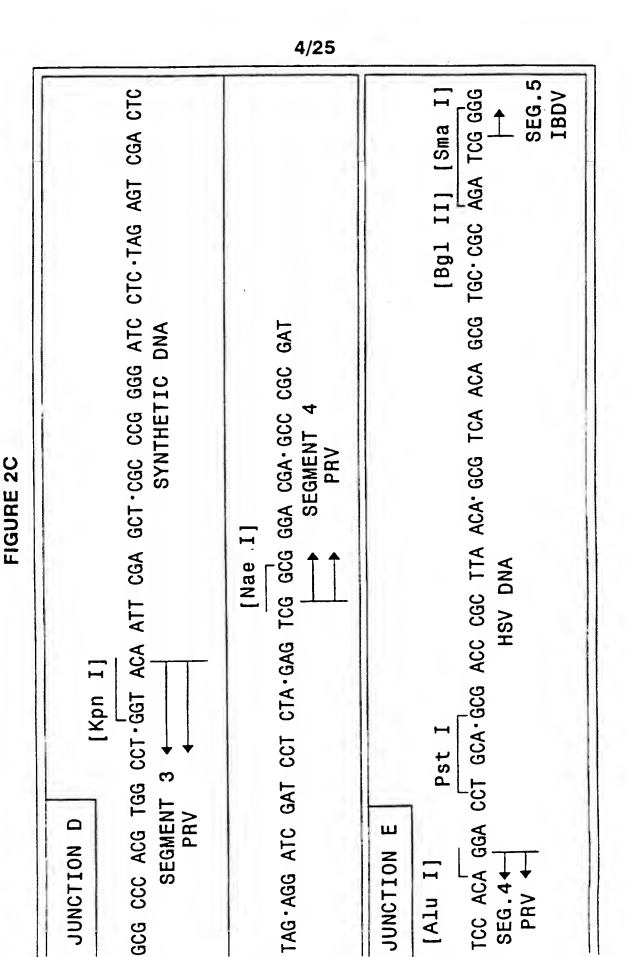
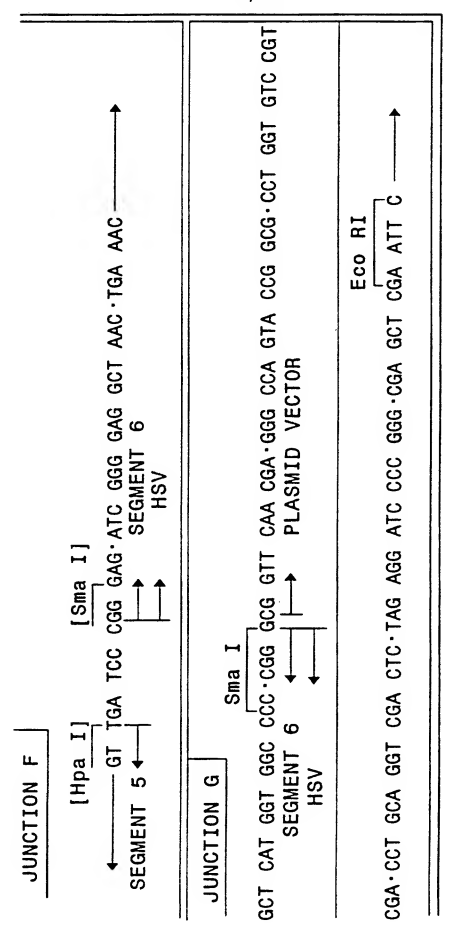


FIGURE 2D





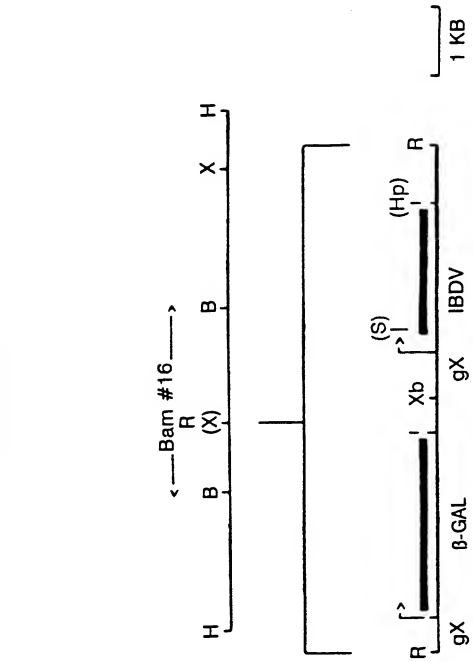
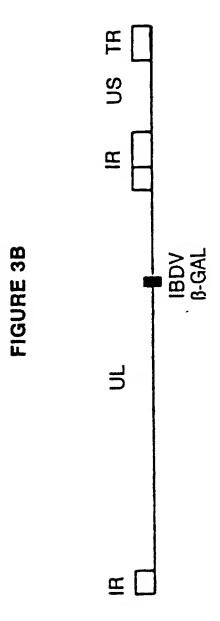


FIGURE 3A



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FIGURE 4

1 2 3 4 5 6 7 kDa

97.4

43.0

18.4

68.0

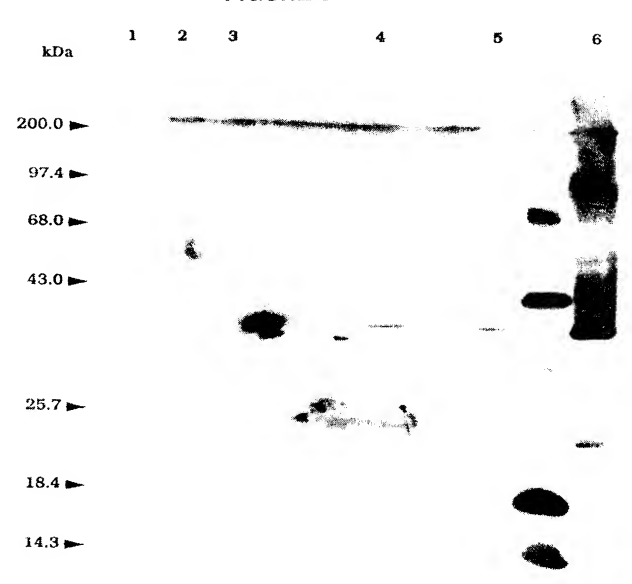
•

**f** 

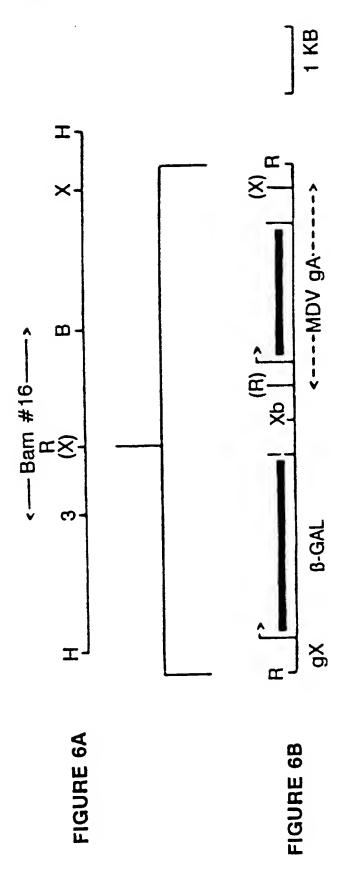
25.7

14.3

## FIGURE 5







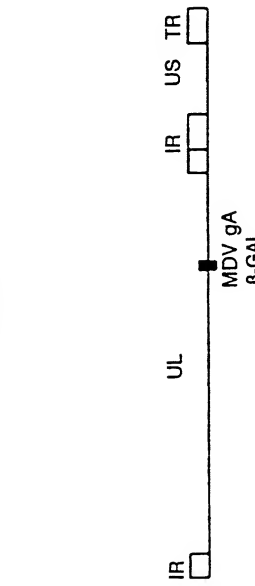
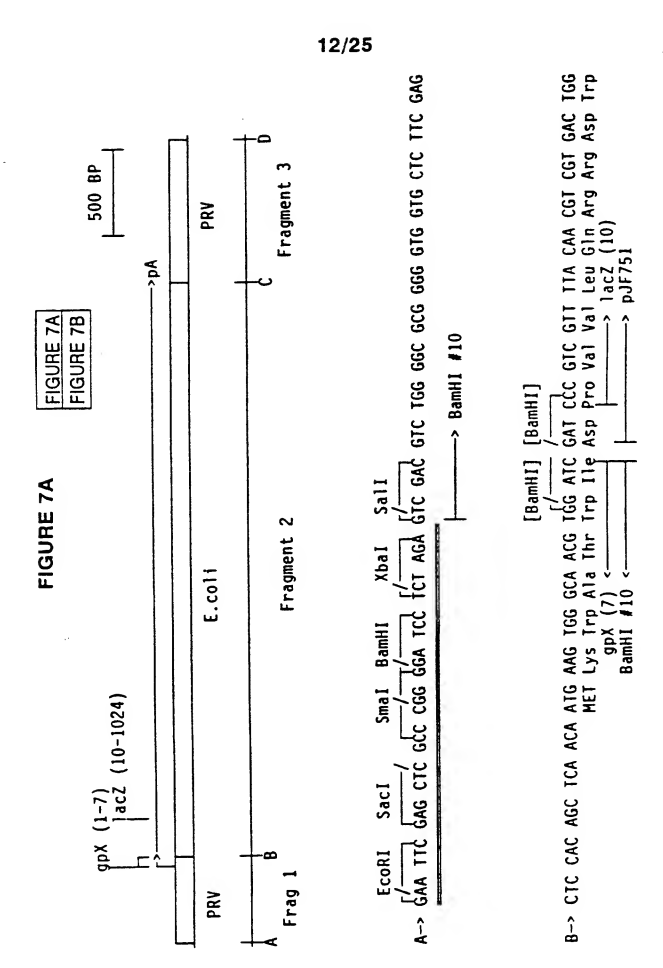
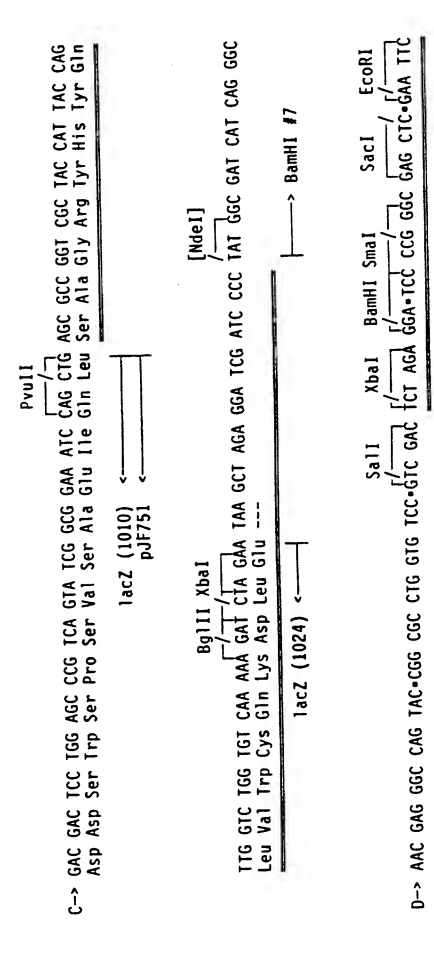


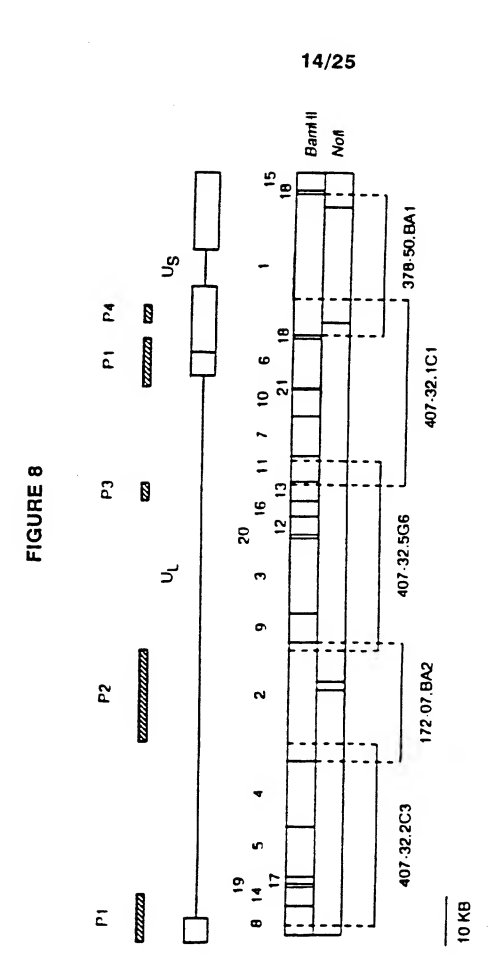
FIGURE 6C

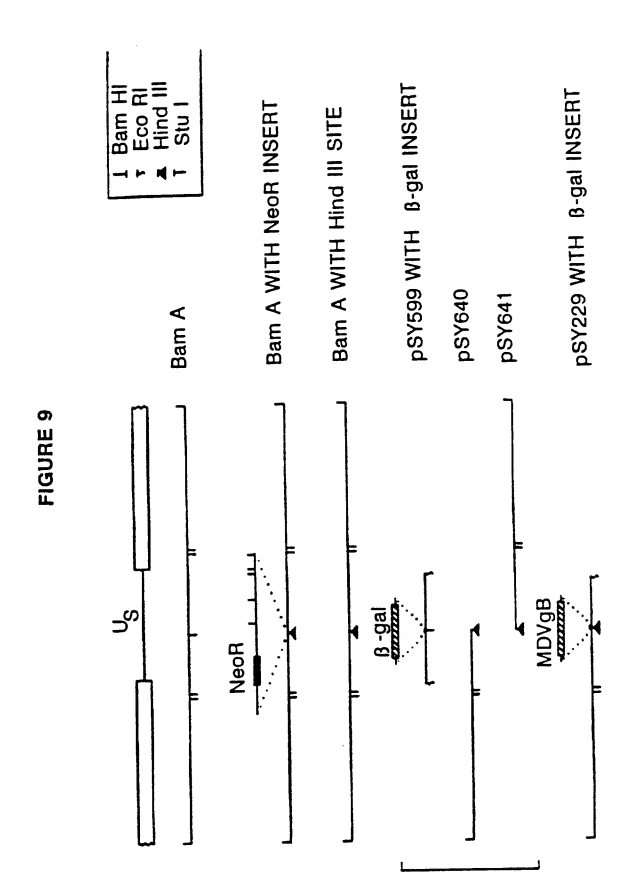


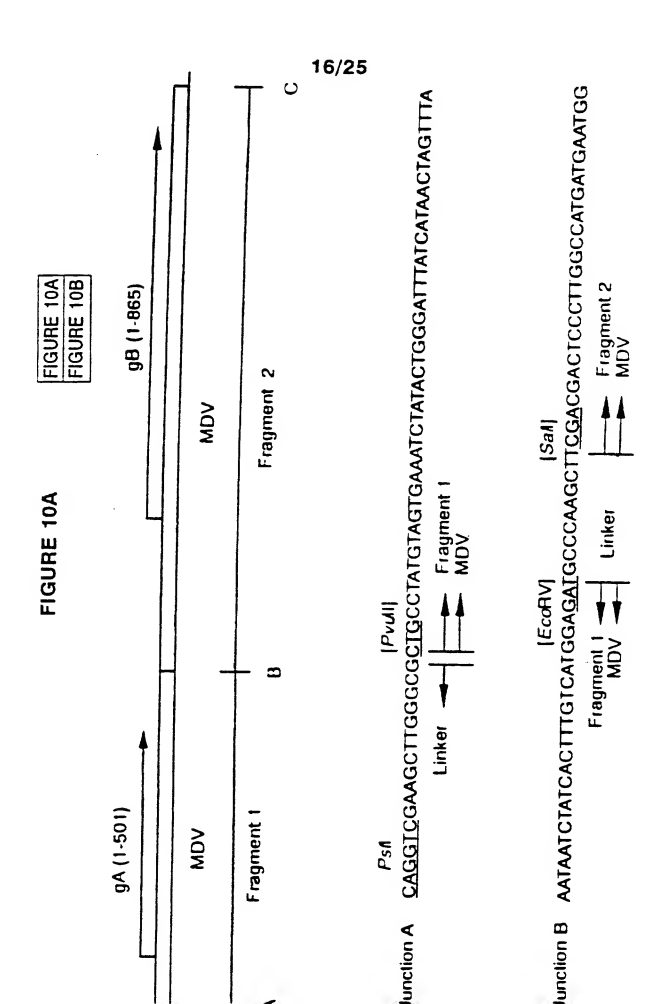
## FIGURE 7B



BamHI #7







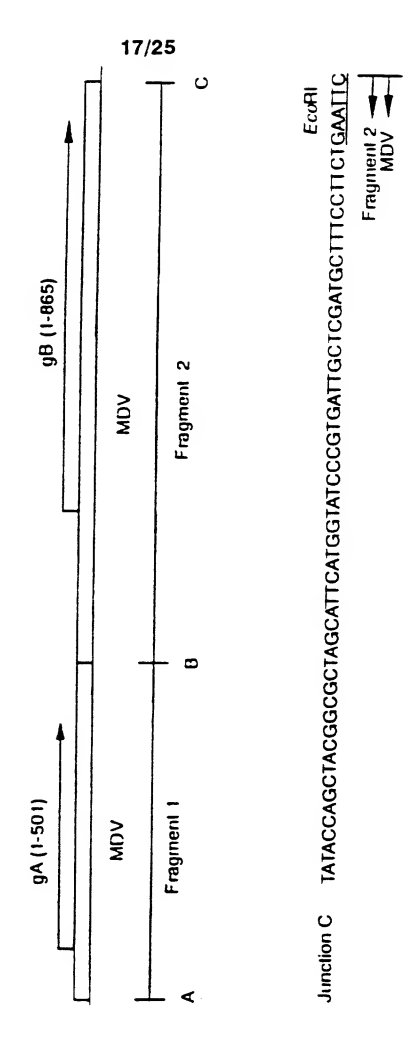
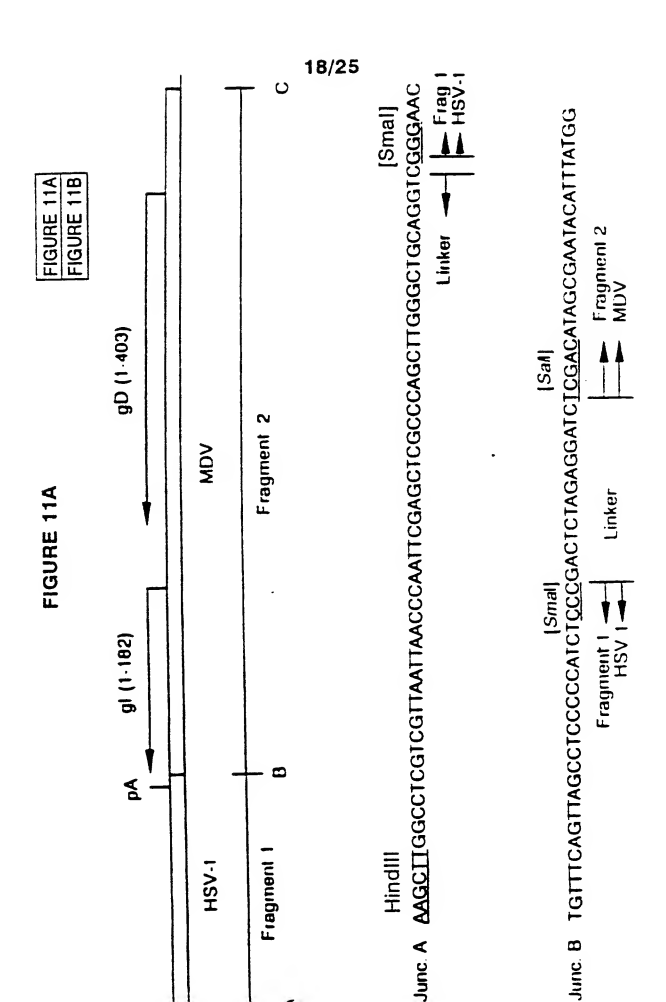


FIGURE 10B



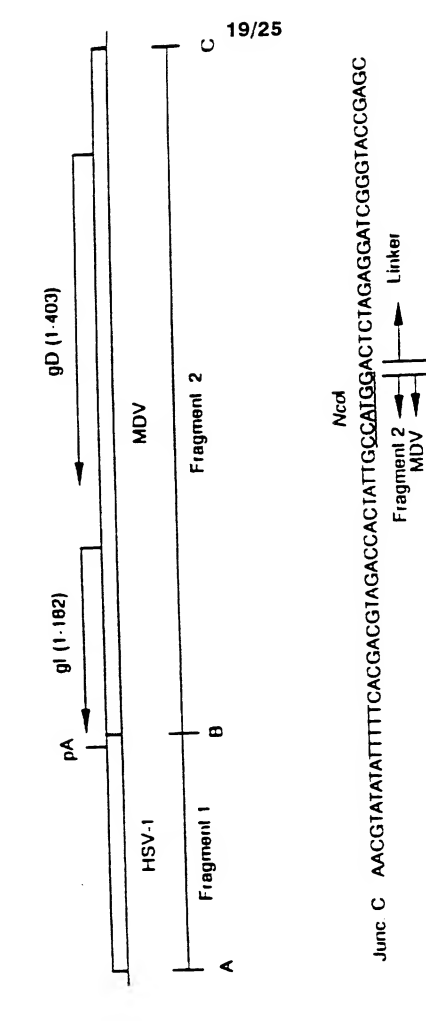
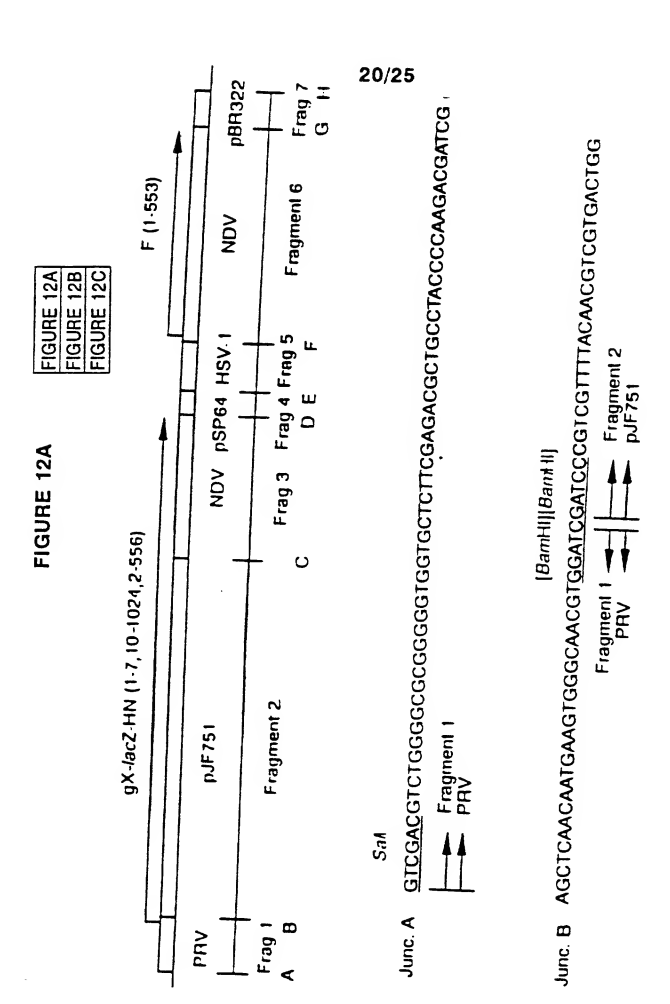


FIGURE 11B

June C TCGAATTGGGAAGCTTGTCGACTTAATTAAGCGGCCGCGTTTAAACGGCCCTCGAGGCCAAGCTT

Hindill



# FIGURE 12B

Junc. C GAGCCCGTCAGTATCGGCGGAAATCCAGCTGAGCGCCGGGTCGCTACCATTACCAGTTGGT **→** Linker Pvdl Fragment 2 -

Avall

GTTGGTCTGGTGTCAAAAAGATCC<u>GGACC</u>GCGCCGTTAGCCAAGTTGCGTTAGAGAATGA Fragment 3 NDV Linker + Junc C cont

ACACAGT CACACT CAT GGGGGCCGAAGGCAGAATT CGTAAT CAT GGT CATAGCT GTTT CC Fragment 4 pSP64 EcoRI Fragment 3 -Junc. D

AAACCTGTCGTGC<u>CAG</u>CGAGCTCGGGATCCTCTAGAGGATC<u>CCCGGGG</u>CCCCGCCCCTGC Smal [Pvd] Junc E

Fragment 4 pSP64

Fragment 5
HSV.1

## FIGURE 12C

Junc. F TCGTCCACACGGAGCGCGGCTGCCGACACGGAICCCGGTTGGCGCCCCTCCAGGTGCAGGA Bantill

Fragment 5 A HSV.1

Fragment 6 NDV

22/25 June G AACCCCCCCCCCCCCCCCCCCCCCCCCCGIGCAGCCATCGIGGIGICACGCTCGICGTTIG PsA

Fragment 7 pBR322 

Junc. H TGTCATGCCATCCGTAAGATGCTTTTCTGTGACTGGTGAGICGGATCCTCTAGAGICGAC - Linker [Scal]

Fragment 7 - pBR322 -

FIGURE 13A

24/25

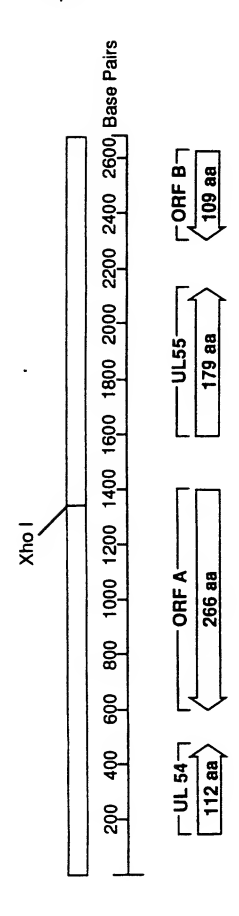
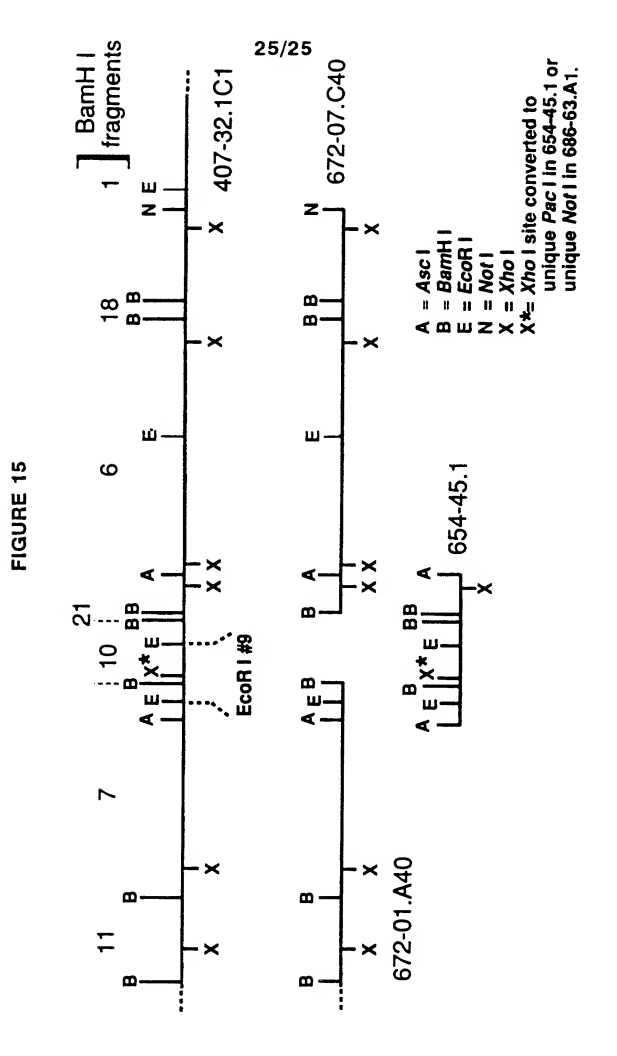


FIGURE 12



## INTERNATIONAL SEARCH REPORT

Form PCT/ISA/210 (second sheet)(July 1992)\*

Inter. onal application No. PCT/US95/10245

A. CLASSIFICATION OF SUBJECT MATTER		
IPC(6) :Please See Extra Sheet. US CL :Please See Extra Sheet.		
According to International Patent Classification (IPC) or to bot	h national classification and IPC	
B. FIELDS SEARCHED	······································	
Minimum documentation searched (classification system follows	ed by classification symbols)	
U.S.: Please See Extra Sheet.		
Documentation searched other than minimum documentation to the	ne extent that such documents are included	in the fields searched
Electronic data base consulted during the international search (n	same of data base and where practicable	search terms used)
APS, Medline, CABA, Agricola, Derwent WPIDS, Inpado vaccine		
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
A US, A, 5,187,087 (SONDERMEI 1993, see entire document	JER ET AL.) 16 February	1-40
A WO 93/25665 (SYNTRO CORPO 1993, SEE ENTIRE DOCUMENT	WO 93/25665 (SYNTRO CORPORATION) 23 DECEMBER 1993, SEE ENTIRE DOCUMENT	
A Vaccine, Volume 11, Number 3, i et al., "Avian herpesvirus as a expression of heterologous antig entire document	live viral vector for the	1-40
X Further documents are listed in the continuation of Box (	See patent family annex.	
Special categories of cited documents:     T later document published after the international filing date or priority		
"A" document defining the general state of the art which is not considered	date and not in conflict with the application but cred to understand the document defining the general state of the art which is not considered principle or theory underlying the invention	
to be of particular relevance  E' cartier document published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be	
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other	considered novel or cannot be considered to involve an inventive step when the document is taken alone  "Y" document of particular relevance; the claimed invention cannot be	
"O" document referring to an oral disclosure, use, exhibition or other means	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art	
*P* document published prior to the international filing date but later than the priority date claimed	*&* document member of the same patent family	
Date of the actual completion of the international search  Date of mailing of the international search report		
28 OCTOBER 1995 28 NOV 1995		
Commissioner of Patents and Trademarks  Box PCT  Authorized officer  LAWRENCE I CARROLL II		
Washington, D.C. 20231 acsimile No. (703) 305-3230 Telephone No. (703) 308-0196		

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
•	Journal of General Virology, Volume 74, issued 1993, Ross et al., "Construction and properties of a turkey herpesvirus recombinant expressing the Marek's disease virus homologue of glycoprotein B of herpes simplex virus", pages 371-377, see entire document	1-40
	Proceedings of the National Academy of Sciences, Volume 89, issued April 1992, "Vaccinia virus recombinants expressing chimeric proteins of human immunodeficiency virus and gamma interferon are attenuated for nude mice", pages 3409-3413, see abstract	1-40
	•	

## INTERNATIONAL SEARCH REPORT

Inter .onal application No. PCT/US95/10245

### A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6):

C12N 5/10, 5/20, 7/01, 15/00, 15/09, 15/12, 15/19, 15/24, 15/26, 15/27, 15/34, 15/38, 15/40, 15/45, 15/86; A61K 39/12, 39/295, 39/17, 39/245, 39/255, 39/265, 39/215

## A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

424/186.1, 199.1, 201.1, 202.1, 204.1, 214.1, 229.1, 222.1; 435/69.3, 69.1, 235.1, 240.1, 240.2 320.1; 536/23.72, 24.2, 23.51, 23.52, 23.2

### **B. FIELDS SEARCHED**

Minimum documentation searched Classification System: U.S.

424/186.1, 199.1, 201.1, 202.1, 204.1, 214.1, 229.1, 222.1; 435/69.3, 69.1, 235.1, 240.1, 240.2, 320.1; 536/23.72, 24.2, 23.51, 23.52, 23.2